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(54) Title: NOVEL NUCLEIC ACIDS AND POLYPEPTIDES

(57) Abstract: The present invention provides novel nucleic acids, novel polypeptide sequences encoded by these nucleic acids and uses thereof.

#### NOVEL NUCLEIC ACIDS AND POLYPEPTIDES

#### 1. TECHNICAL FIELD

The present invention provides novel polynucleotides and proteins encoded by such polynucleotides, along with uses for these polynucleotides and proteins, for example in therapeutic, diagnostic and research methods.

# 2. BACKGROUND

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Technology aimed at the discovery of protein factors (including e.g., cytokines, such as lymphokines, interferons, CSFs, chemokines, and interleukins) has matured rapidly over the past decade. The now routine hybridization cloning and expression cloning techniques clone novel polynucleotides "directly" in the sense that they rely on information directly related to the discovered protein (i.e., partial DNA/amino acid sequence of the protein in the case of hybridization cloning; activity of the protein in the case of expression cloning). More recent "indirect" cloning techniques such as signal sequence cloning, which isolates DNA sequences based on the presence of a now well-recognized secretory leader sequence motif, as well as various PCR-based or low stringency hybridization-based cloning techniques, have advanced the state of the art by making available large numbers of DNA/amino acid sequences for proteins that are known to have biological activity, for example, by virtue of their secreted nature in the case of leader sequence cloning, by virtue of their cell or tissue source in the case of PCR-based techniques, or by virtue of structural similarity to other genes of known biological activity.

Identified polynucleotide and polypeptide sequences have numerous applications in, for example, diagnostics, forensics, gene mapping; identification of mutations responsible for genetic disorders or other traits, to assess biodiversity, and to produce many other types of data and products dependent on DNA and amino acid sequences.

# 3. SUMMARY OF THE INVENTION

The compositions of the present invention include novel isolated polypeptides, novel isolated polynucleotides encoding such polypeptides, including recombinant DNA molecules, cloned genes or degenerate variants thereof, especially naturally occurring variants such as allelic variants, antisense polynucleotide molecules, and antibodies that specifically recognize one or more epitopes present on such polypeptides, as well as hybridomas producing such antibodies.

The compositions of the present invention additionally include vectors, including expression vectors, containing the polynucleotides of the invention, cells genetically engineered to contain such polynucleotides and cells genetically engineered to express such polynucleotides.

The present invention relates to a collection or library of at least one novel nucleic acid sequence assembled from expressed sequence tags (ESTs) isolated mainly by sequencing by hybridization (SBH), and in some cases, sequences obtained from one or more public databases. The invention relates also to the proteins encoded by such polynucleotides, along with therapeutic, diagnostic and research utilities for these polynucleotides and proteins. These nucleic acid sequences are designated as SEQ ID NO: 1-1350. The polypeptides sequences are designated SEQ ID NO: 1351-2700. The nucleic acids and polypeptides are provided in the Sequence Listing. In the nucleic acids provided in the Sequence Listing, A is adenosine; C is cytosine; G is guanine; T is thymine; and N is any of the four bases. In the amino acids provided in the Sequence Listing, \* corresponds to the stop codon.

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The nucleic acid sequences of the present invention also include, nucleic acid sequences that hybridize to the complement of SEQ ID NO:1-1350 under stringent hybridization conditions; nucleic acid sequences which are allelic variants or species homologues of any of the nucleic acid sequences recited above, or nucleic acid sequences that encode a peptide comprising a specific domain or truncation of the peptides encoded by SEQ ID NO:1-1350. A polynucleotide comprising a nucleotide sequence having at least 90% identity to an identifying sequence of SEQ ID NO:1-1350 or a degenerate variant or fragment thereof. The identifying sequence can be 100 base pairs in length.

The nucleic acid sequences of the present invention also include the sequence information from the nucleic acid sequences of SEQ ID NO:1-1350. The sequence information can be a segment of any one of SEQ ID NO:1-1350 that uniquely identifies or represents the sequence information of SEQ ID NO:1-1350.

A collection as used in this application can be a collection of only one polynucleotide. The collection of sequence information or identifying information of each sequence can be provided on a nucleic acid array. In one embodiment, segments of sequence information is provided on a nucleic acid array to detect the polynucleotide that contains the segment. The array can be designed to detect full-match or mismatch to the polynucleotide that contains the segment. The collection can also be provided in a computer-readable format.

This invention also includes the reverse or direct complement of any of the nucleic acid sequences recited above; cloning or expression vectors containing the nucleic acid sequences; and host cells or organisms transformed with these expression vectors. Nucleic acid sequences (or their reverse or direct complements) according to the invention have numerous applications in a variety of techniques known to those skilled in the art of molecular biology, such as use as hybridization probes, use as primers for PCR, use in an array, use in computer-readable media, use in sequencing

full-length genes, use for chromosome and gene mapping, use in the recombinant production of protein, and use in the generation of anti-sense DNA or RNA, their chemical analogs and the like.

In a preferred embodiment, the nucleic acid sequences of SEQ ID NO:1-1350 or novel segments or parts of the nucleic acids of the invention are used as primers in expression assays that are well known in the art. In a particularly preferred embodiment, the nucleic acid sequences of SEQ ID NO:1-1350 or novel segments or parts of the nucleic acids provided herein are used in diagnostics for identifying expressed genes or, as well known in the art and exemplified by Vollrath et al., Science 258:52-59 (1992), as expressed sequence tags for physical mapping of the human genome.

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The isolated polynucleotides of the invention include, but are not limited to, a polynucleotide comprising any one of the nucleotide sequences set forth in SEQ ID NO:1-1350; a polynucleotide comprising any of the full length protein coding sequences of SEQ ID NO:1 - 1350; and a polynucleotide comprising any of the nucleotide sequences of the mature protein coding sequences of SEQ ID NO: 1-1350. The polynucleotides of the present invention also include, but are not limited to, a polynucleotide that hybridizes under stringent hybridization conditions to (a) the complement of any one of the nucleotide sequences set forth in SEQ ID NO:1-1350; (b) a nucleotide sequence encoding any one of the amino acid sequences set forth in the Sequence Listing (e.g., SEQ ID NO: 1351-2700); (c) a polynucleotide which is an allelic variant of any polynucleotides recited above; (d) a polynucleotide which encodes a species homolog (e.g. orthologs) of any of the proteins recited above; or (e) a polynucleotide that encodes a polypeptide comprising a specific domain or truncation of any of the polypeptides comprising an amino acid sequence set forth in the Sequence Listing.

The isolated polypeptides of the invention include, but are not limited to, a polypeptide comprising any of the amino acid sequences set forth in the Sequence Listing; or the corresponding full length or mature protein. Polypeptides of the invention also include polypeptides with biological activity that are encoded by (a) any of the polynucleotides having a nucleotide sequence set forth in SEQ ID NO:1-1350; or (b) polynucleotides that hybridize to the complement of the polynucleotides of (a) under stringent hybridization conditions. Biologically or immunologically active variants of any of the polypeptide sequences in the Sequence Listing, and "substantial equivalents" thereof (e.g., with at least about 65%, 70%, 75%, 80%, 85%, 90%, 95%, 98% or 99% amino acid sequence identity) that preferably retain biological activity are also contemplated. The polypeptides of the invention may be wholly or partially chemically synthesized but are preferably produced by recombinant means using the genetically engineered cells (e.g. host cells) of the invention.

The invention also provides compositions comprising a polypeptide of the invention. Polypeptide compositions of the invention may further comprise an acceptable carrier, such as a hydrophilic, *e.g.*, pharmaceutically acceptable, carrier.

The invention also provides host cells transformed or transfected with a polynucleotide of the invention.

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The invention also relates to methods for producing a polypeptide of the invention comprising growing a culture of the host cells of the invention in a suitable culture medium under conditions permitting expression of the desired polypeptide, and purifying the polypeptide from the culture or from the host cells. Preferred embodiments include those in which the protein produced by such process is a mature form of the protein.

Polynucleotides according to the invention have numerous applications in a variety of techniques known to those skilled in the art of molecular biology. These techniques include use as hybridization probes, use as oligomers, or primers, for PCR, use for chromosome and gene mapping, use in the recombinant production of protein, and use in generation of anti-sense DNA or RNA, their chemical analogs and the like. For example, when the expression of an mRNA is largely restricted to a particular cell or tissue type, polynucleotides of the invention can be used as hybridization probes to detect the presence of the particular cell or tissue mRNA in a sample using, *e.g.*, *in situ* hybridization.

In other exemplary embodiments, the polynucleotides are used in diagnostics as expressed sequence tags for identifying expressed genes or, as well known in the art and exemplified by Vollrath et al., Science 258:52-59 (1992), as expressed sequence tags for physical mapping of the human genome.

The polypeptides according to the invention can be used in a variety of conventional procedures and methods that are currently applied to other proteins. For example, a polypeptide of the invention can be used to generate an antibody that specifically binds the polypeptide. Such antibodies, particularly monoclonal antibodies, are useful for detecting or quantitating the polypeptide in tissue. The polypeptides of the invention can also be used as molecular weight markers, and as a food supplement.

Methods are also provided for preventing, treating, or ameliorating a medical condition which comprises the step of administering to a mammalian subject a therapeutically effective amount of a composition comprising a polypeptide of the present invention and a pharmaceutically acceptable carrier.

In particular, the polypeptides and polynucleotides of the invention can be utilized, for example, in methods for the prevention and/or treatment of disorders involving aberrant protein expression or biological activity.

The present invention further relates to methods for detecting the presence of the polynucleotides or polypeptides of the invention in a sample. Such methods can, for example, be utilized as part of prognostic and diagnostic evaluation of disorders as recited herein and for the identification of subjects exhibiting a predisposition to such conditions. The invention provides a method for detecting the polynucleotides of the invention in a sample, comprising contacting the sample with a compound that binds to and forms a complex with the polynucleotide of interest for a period sufficient to form the complex and under conditions sufficient to form a complex and detecting the complex such that if a complex is detected, the polynucleotide of interest is detected. The invention also provides a method for detecting the polypeptides of the invention in a sample comprising contacting the sample with a compound that binds to and form a complex with the polypeptide under conditions and for a period sufficient to form the complex and detecting the formation of the complex such that if a complex is formed, the polypeptide is detected.

The invention also provides kits comprising polynucleotide probes and/or monoclonal antibodies, and optionally quantitative standards, for carrying out methods of the invention. Furthermore, the invention provides methods for evaluating the efficacy of drugs, and monitoring the progress of patients, involved in clinical trials for the treatment of disorders as recited above.

The invention also provides methods for the identification of compounds that modulate (i.e., increase or decrease) the expression or activity of the polynucleotides and/or polypeptides of the invention. Such methods can be utilized, for example, for the identification of compounds that can ameliorate symptoms of disorders as recited herein. Such methods can include, but are not limited to, assays for identifying compounds and other substances that interact with (e.g., bind to) the polypeptides of the invention. The invention provides a method for identifying a compound that binds to the polypeptides of the invention comprising contacting the compound with a polypeptide of the invention in a cell for a time sufficient to form a polypeptide/compound complex, wherein the complex drives expression of a reporter gene sequence in the cell; and detecting the complex by detecting the reporter gene sequence expression such that if expression of the reporter gene is detected the compound the binds to a polypeptide of the invention is identified.

The methods of the invention also provides methods for treatment which involve the administration of the polynucleotides or polypeptides of the invention to individuals exhibiting symptoms or tendencies. In addition, the invention encompasses methods for treating diseases o disorders as recited herein comprising administering compounds and other substances that modulate the overall activity of the target gene products. Compounds and other substances can

effect such modulation either on the level of target gene/protein expression or target protein activity.

The polypeptides of the present invention and the polynucleotides encoding them are also useful for the same functions known to one of skill in the art as the polypeptides and polynucleotides to which they have homology (set forth in Table 2). If no homology is set forth for a sequence, then the polypeptides and polynucleotides of the present invention are useful for a variety of applications, as described herein, including use in arrays for detection.

#### 4. DETAILED DESCRIPTION OF THE INVENTION

#### 4.1 DEFINITIONS

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It must be noted that as used herein and in the appended claims, the singular forms "a", "an" and "the" include plural references unless the context clearly dictates otherwise.

The term "active" refers to those forms of the polypeptide which retain the biologic and/or immunologic activities of any naturally occurring polypeptide. According to the invention, the terms "biologically active" or "biological activity" refer to a protein or peptide having structural, regulatory or biochemical functions of a naturally occurring molecule. Likewise "immunologically active" or "immunological activity" refers to the capability of the natural, recombinant or synthetic polypeptide to induce a specific immune response in appropriate animals or cells and to bind with specific antibodies.

The term "activated cells" as used in this application are those cells which are engaged in extracellular or intracellular membrane trafficking, including the export of secretory or enzymatic molecules as part of a normal or disease process.

The terms "complementary" or "complementarity" refer to the natural binding of polynucleotides by base pairing. For example, the sequence 5'-AGT-3' binds to the complementary sequence 3'-TCA-5'. Complementarity between two single-stranded molecules may be "partial" such that only some of the nucleic acids bind or it may be "complete" such that total complementarity exists between the single stranded molecules. The degree of complementarity between the nucleic acid strands has significant effects on the efficiency and strength of the hybridization between the nucleic acid strands.

The term "embryonic stem cells (ES)" refers to a cell that can give rise to many differentiated cell types in an embryo or an adult, including the germ cells. The term "germ line stem cells (GSCs)" refers to stem cells derived from primordial stem cells that provide a steady and continuous source of germ cells for the production of gametes. The term "primordial germ

cells (PGCs)" refers to a small population of cells set aside from other cell lineages particularly from the yolk sac, mesenteries, or gonadal ridges during embryogenesis that have the potential to differentiate into germ cells and other cells. PGCs are the source from which GSCs and ES cells are derived. The PGCs, the GSCs and the ES cells are capable of self-renewal. Thus these cells not only populate the germ line and give rise to a plurality of terminally differentiated cells that comprise the adult specialized organs, but are able to regenerate themselves.

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The term "expression modulating fragment," EMF, means a series of nucleotides which modulates the expression of an operably linked ORF or another EMF.

As used herein, a sequence is said to "modulate the expression of an operably linked sequence" when the expression of the sequence is altered by the presence of the EMF. EMFs include, but are not limited to, promoters, and promoter modulating sequences (inducible elements). One class of EMFs are nucleic acid fragments which induce the expression of an operably linked ORF in response to a specific regulatory factor or physiological event.

The terms "nucleotide sequence" or "nucleic acid" or "polynucleotide" or "oligonculeotide" are used interchangeably and refer to a heteropolymer of nucleotides or the sequence of these nucleotides. These phrases also refer to DNA or RNA of genomic or synthetic origin which may be single-stranded or double-stranded and may represent the sense or the antisense strand, to peptide nucleic acid (PNA) or to any DNA-like or RNA-like material. In the sequences herein A is adenine, C is cytosine, T is thymine, G is guanine and N is A, C, G or T (U). It is contemplated that where the polynucleotide is RNA, the T (thymine) in the sequences provided herein is substituted with U (uracil). Generally, nucleic acid segments provided by this invention may be assembled from fragments of the genome and short oligonucleotide linkers, or from a series of oligonucleotides, or from individual nucleotides, to provide a synthetic nucleic acid which is capable of being expressed in a recombinant transcriptional unit comprising regulatory elements derived from a microbial or viral operon, or a eukaryotic gene.

The terms "oligonucleotide fragment" or a "polynucleotide fragment", "portion," or "segment" or "probe" or "primer" are used interchangeably and refer to a sequence of nucleotide residues which are at least about 5 nucleotides, more preferably at least about 7 nucleotides, more preferably at least about 11 nucleotides and most preferably at least about 17 nucleotides. The fragment is preferably less than about 500 nucleotides, preferably less than about 200 nucleotides, more preferably less than about 100 nucleotides, more preferably less than about 50 nucleotides and most preferably less than 30 nucleotides. Preferably the probe is from about 6 nucleotides to about 200 nucleotides, preferably from about 15 to about 50 nucleotides, more preferably from about 17 to 30 nucleotides and most preferably from about 20 to 25 nucleotides. Preferably the fragments can

be used in polymerase chain reaction (PCR), various hybridization procedures or microarray procedures to identify or amplify identical or related parts of mRNA or DNA molecules. A fragment or segment may uniquely identify each polynucleotide sequence of the present invention. Preferably the fragment comprises a sequence substantially similar to any one of SEQ ID NOs:1-1350.

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Probes may, for example, be used to determine whether specific mRNA molecules are present in a cell or tissue or to isolate similar nucleic acid sequences from chromosomal DNA as described by Walsh et al. (Walsh, P.S. et al., 1992, PCR Methods Appl 1:241-250). They may be labeled by nick translation, Klenow fill-in reaction, PCR, or other methods well known in the art. Probes of the present invention, their preparation and/or labeling are elaborated in Sambrook, J. et al., 1989, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, NY; or Ausubel, F.M. et al., 1989, Current Protocols in Molecular Biology, John Wiley & Sons, New York NY, both of which are incorporated herein by reference in their entirety.

The nucleic acid sequences of the present invention also include the sequence information from the nucleic acid sequences of SEQ ID NO:1-1350. The sequence information can be a segment of any one of SEQ ID NO:1-1350 that uniquely identifies or represents the sequence information of that sequence of SEQ ID NO:1-1350. One such segment can be a twenty-mer nucleic acid sequence because the probability that a twenty-mer is fully matched in the human genome is 1 in 300. In the human genome, there are three billion base pairs in one set of chromosomes. Because 4<sup>20</sup> possible twenty-mers exist, there are 300 times more twenty-mers than there are base pairs in a set of human chromosomes. Using the same analysis, the probability for a seventeen-mer to be fully matched in the human genome is approximately 1 in 5. When these segments are used in arrays for expression studies, fifteen-mer segments can be used. The probability that the fifteen-mer is fully matched in the expressed sequences is also approximately one in five because expressed sequences comprise less than approximately 5% of the entire genome sequence.

Similarly, when using sequence information for detecting a single mismatch, a segment can be a twenty-five mer. The probability that the twenty-five mer would appear in a human genome with a single mismatch is calculated by multiplying the probability for a full match  $(1 \div 4^{25})$  times the increased probability for mismatch at each nucleotide position  $(3 \times 25)$ . The probability that an eighteen mer with a single mismatch can be detected in an array for expression studies is approximately one in five. The probability that a twenty-mer with a single mismatch can be detected in a human genome is approximately one in five.

The term "open reading frame," ORF, means a series of nucleotide triplets coding for amino acids without any termination codons and is a sequence translatable into protein.

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The terms "operably linked" or "operably associated" refer to functionally related nucleic acid sequences. For example, a promoter is operably associated or operably linked with a coding sequence if the promoter controls the transcription of the coding sequence. While operably linked nucleic acid sequences can be contiguous and in the same reading frame, certain genetic elements *e.g.* repressor genes are not contiguously linked to the coding sequence but still control transcription/translation of the coding sequence.

The term "pluripotent" refers to the capability of a cell to differentiate into a number of differentiated cell types that are present in an adult organism. A pluripotent cell is restricted in its differentiation capability in comparison to a totipotent cell.

The terms "polypeptide" or "peptide" or "amino acid sequence" refer to an oligopeptide, peptide, polypeptide or protein sequence or fragment thereof and to naturally occurring or synthetic molecules. A polypeptide "fragment," "portion," or "segment" is a stretch of amino acid residues of at least about 5 amino acids, preferably at least about 7 amino acids, more preferably at least about 9 amino acids and most preferably at least about 17 or more amino acids. The peptide preferably is not greater than about 200 amino acids, more preferably less than 150 amino acids and most preferably less than 100 amino acids. Preferably the peptide is from about 5 to about 200 amino acids. To be active, any polypeptide must have sufficient length to display biological and/or immunological activity.

The term "naturally occurring polypeptide" refers to polypeptides produced by cells that have not been genetically engineered and specifically contemplates various polypeptides arising from post-translational modifications of the polypeptide including, but not limited to, acetylation, carboxylation, glycosylation, phosphorylation, lipidation and acylation.

The term "translated protein coding portion" means a sequence which encodes for the full length protein which may include any leader sequence or any processing sequence.

The term "mature protein coding sequence" means a sequence which encodes a peptide or protein without a signal or leader sequence. The "mature protein portion" means that portion of the protein which does not include a signal or leader sequence. The peptide may have been produced by processing in the cell which removes any leader/signal sequence. The mature protein portion may or may not include the initial methionine residue. The methionine residue may be removed from the protein during processing in the cell. The peptide may be produced synthetically or the protein may have been produced using a polynucleotide only encoding for the mature protein coding sequence.

The term "derivative" refers to polypeptides chemically modified by such techniques as ubiquitination, labeling (e.g., with radionuclides or various enzymes), covalent polymer attachment such as pegylation (derivatization with polyethylene glycol) and insertion or substitution by chemical synthesis of amino acids such as ornithine, which do not normally occur in human proteins.

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The term "variant" (or "analog") refers to any polypeptide differing from naturally occurring polypeptides by amino acid insertions, deletions, and substitutions, created using, e g., recombinant DNA techniques. Guidance in determining which amino acid residues may be replaced, added or deleted without abolishing activities of interest, may be found by comparing the sequence of the particular polypeptide with that of homologous peptides and minimizing the number of amino acid sequence changes made in regions of high homology (conserved regions) or by replacing amino acids with consensus sequence.

Alternatively, recombinant variants encoding these same or similar polypeptides may be synthesized or selected by making use of the "redundancy" in the genetic code. Various codon substitutions, such as the silent changes which produce various restriction sites, may be introduced to optimize cloning into a plasmid or viral vector or expression in a particular prokaryotic or eukaryotic system. Mutations in the polynucleotide sequence may be reflected in the polypeptide or domains of other peptides added to the polypeptide to modify the properties of any part of the polypeptide, to change characteristics such as ligand-binding affinities, interchain affinities, or degradation/turnover rate.

Preferably, amino acid "substitutions" are the result of replacing one amino acid with another amino acid having similar structural and/or chemical properties, *i.e.*, conservative amino acid replacements. "Conservative" amino acid substitutions may be made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophilicity, and/or the amphipathic nature of the residues involved. For example, nonpolar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan, and methionine; polar neutral amino acids include glycine, serine, threonine, cysteine, tyrosine, asparagine, and glutamine; positively charged (basic) amino acids include arginine, lysine, and histidine; and negatively charged (acidic) amino acids include aspartic acid and glutamic acid. "Insertions" or "deletions" are preferably in the range of about 1 to 20 amino acids, more preferably 1 to 10 amino acids. The variation allowed may be experimentally determined by systematically making insertions, deletions, or substitutions of amino acids in a polypeptide molecule using recombinant DNA techniques and assaying the resulting recombinant variants for activity.

Alternatively, where alteration of function is desired, insertions, deletions or non-conservative alterations can be engineered to produce altered polypeptides. Such alterations

can, for example, alter one or more of the biological functions or biochemical characteristics of the polypeptides of the invention. For example, such alterations may change polypeptide characteristics such as ligand-binding affinities, interchain affinities, or degradation/turnover rate. Further, such alterations can be selected so as to generate polypeptides that are better suited for expression, scale up and the like in the host cells chosen for expression. For example, cysteine residues can be deleted or substituted with another amino acid residue in order to eliminate disulfide bridges.

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The terms "purified" or "substantially purified" as used herein denotes that the indicated nucleic acid or polypeptide is present in the substantial absence of other biological macromolecules, *e.g.*, polynucleotides, proteins, and the like. In one embodiment, the polynucleotide or polypeptide is purified such that it constitutes at least 95% by weight, more preferably at least 99% by weight, of the indicated biological macromolecules present (but water, buffers, and other small molecules, especially molecules having a molecular weight of less than 1000 daltons, can be present).

The term "isolated" as used herein refers to a nucleic acid or polypeptide separated from at least one other component (e.g., nucleic acid or polypeptide) present with the nucleic acid or polypeptide in its natural source. In one embodiment, the nucleic acid or polypeptide is found in the presence of (if anything) only a solvent, buffer, ion, or other component normally present in a solution of the same. The terms "isolated" and "purified" do not encompass nucleic acids or polypeptides present in their natural source.

The term "recombinant," when used herein to refer to a polypeptide or protein, means that a polypeptide or protein is derived from recombinant (e.g., microbial, insect, or mammalian) expression systems. "Microbial" refers to recombinant polypeptides or proteins made in bacterial or fungal (e.g., yeast) expression systems. As a product, "recombinant microbial" defines a polypeptide or protein essentially free of native endogenous substances and unaccompanied by associated native glycosylation. Polypeptides or proteins expressed in most bacterial cultures, e.g., E. coli, will be free of glycosylation modifications; polypeptides or proteins expressed in yeast will have a glycosylation pattern in general different from those expressed in mammalian cells.

The term "recombinant expression vehicle or vector" refers to a plasmid or phage or virus or vector, for expressing a polypeptide from a DNA (RNA) sequence. An expression vehicle can comprise a transcriptional unit comprising an assembly of (1) a genetic element or elements having a regulatory role in gene expression, for example, promoters or enhancers, (2) a structural or coding sequence which is transcribed into mRNA and translated into protein, and (3) appropriate transcription initiation and termination sequences. Structural units intended for use

in yeast or eukaryotic expression systems preferably include a leader sequence enabling extracellular secretion of translated protein by a host cell. Alternatively, where recombinant protein is expressed without a leader or transport sequence, it may include an amino terminal methionine residue. This residue may or may not be subsequently cleaved from the expressed recombinant protein to provide a final product.

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The term "recombinant expression system" means host cells which have stably integrated a recombinant transcriptional unit into chromosomal DNA or carry the recombinant transcriptional unit extrachromosomally. Recombinant expression systems as defined herein will express heterologous polypeptides or proteins upon induction of the regulatory elements linked to the DNA segment or synthetic gene to be expressed. This term also means host cells which have stably integrated a recombinant genetic element or elements having a regulatory role in gene expression, for example, promoters or enhancers. Recombinant expression systems as defined herein will express polypeptides or proteins endogenous to the cell upon induction of the regulatory elements linked to the endogenous DNA segment or gene to be expressed. The cells can be prokaryotic or eukaryotic.

The term "secreted" includes a protein that is transported across or through a membrane, including transport as a result of signal sequences in its amino acid sequence when it is expressed in a suitable host cell. "Secreted" proteins include without limitation proteins secreted wholly (e.g., soluble proteins) or partially (e.g., receptors) from the cell in which they are expressed. "Secreted" proteins also include without limitation proteins that are transported across the membrane of the endoplasmic reticulum. "Secreted" proteins are also intended to include proteins containing non-typical signal sequences (e.g. Interleukin-1 Beta, see Krasney, P.A. and Young, P.R. (1992) Cytokine 4(2):134-143) and factors released from damaged cells (e.g. Interleukin-1 Receptor Antagonist, see Arend, W.P. et. al. (1998) Annu. Rev. Immunol. 16:27-55)

Where desired, an expression vector may be designed to contain a "signal or leader sequence" which will direct the polypeptide through the membrane of a cell. Such a sequence may be naturally present on the polypeptides of the present invention or provided from heterologous protein sources by recombinant DNA techniques.

The term "stringent" is used to refer to conditions that are commonly understood in the art as stringent. Stringent conditions can include highly stringent conditions (*i.e.*, hybridization to filter-bound DNA in 0.5 M NaHPO<sub>4</sub>, 7% sodium dodecyl sulfate (SDS), 1 mM EDTA at 65°C, and washing in 0.1X SSC/0.1% SDS at 68°C), and moderately stringent conditions (*i.e.*, washing in 0.2X SSC/0.1% SDS at 42°C). Other exemplary hybridization conditions are described herein in the examples.

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In instances of hybridization of deoxyoligonucleotides, additional exemplary stringent hybridization conditions include washing in 6X SSC/0.05% sodium pyrophosphate at 37°C (for 14-base oligonucleotides), 48°C (for 17-base oligos), 55°C (for 20-base oligonucleotides), and 60°C (for 23-base oligonucleotides).

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As used herein, "substantially equivalent" can refer both to nucleotide and amino acid sequences, for example a mutant sequence, that varies from a reference sequence by one or more substitutions, deletions, or additions, the net effect of which does not result in an adverse functional dissimilarity between the reference and subject sequences. Typically, such a substantially equivalent sequence varies from one of those listed herein by no more than about 35% (i.e., the number of individual residue substitutions, additions, and/or deletions in a substantially equivalent sequence, as compared to the corresponding reference sequence, divided by the total number of residues in the substantially equivalent sequence is about 0.35 or less). Such a sequence is said to have 65% sequence identity to the listed sequence. In one embodiment, a substantially equivalent, e.g., mutant, sequence of the invention varies from a listed sequence by no more than 30% (70% sequence identity); in a variation of this embodiment, by no more than 25% (75% sequence identity); and in a further variation of this embodiment, by no more than 20% (80% sequence identity) and in a further variation of this embodiment, by no more than 10% (90% sequence identity) and in a further variation of this embodiment, by no more that 5% (95% sequence identity). Substantially equivalent, e.g., mutant, amino acid sequences according to the invention preferably have at least 80% sequence identity with a listed amino acid sequence, more preferably at least 85% sequence identity, more preferably at least 90% sequence identity, more preferably at least 95% identity, more preferably at least 98% identity, and most preferably at least 99% identity. Substantially equivalent nucleotide sequences of the invention can have lower percent sequence identities, taking into account, for example, the redundancy or degeneracy of the genetic code. Preferably, nucleotide sequence has at least about 65% identity, more preferably at least about 75% identity, more preferably at least about 80% sequence identity, more preferably at least about 85% sequence identity, more preferably at least about 90% sequence identity, and most preferably at least about 95% identity, more preferably at least about 98% sequence identity, and most preferably at least about 99% sequence identity. For the purposes of the present invention, sequences having substantially equivalent biological activity and substantially equivalent expression characteristics are considered substantially equivalent. For the purposes of determining equivalence, truncation of the mature sequence (e.g., via a mutation which creates a spurious stop codon) should be disregarded. Sequence identity may be determined, e.g., using the Jotun Hein method (Hein, J.

(1990) Methods Enzymol. 183:626-645). Identity between sequences can also be determined by other methods known in the art, e.g. by varying hybridization conditions.

The term "totipotent" refers to the capability of a cell to differentiate into all of the cell types of an adult organism.

The term "transformation" means introducing DNA into a suitable host cell so that the DNA is replicable, either as an extrachromosomal element, or by chromosomal integration. The term "transfection" refers to the taking up of an expression vector by a suitable host cell, whether or not any coding sequences are in fact expressed. The term "infection" refers to the introduction of nucleic acids into a suitable host cell by use of a virus or viral vector.

As used herein, an "uptake modulating fragment," UMF, means a series of nucleotides which mediate the uptake of a linked DNA fragment into a cell. UMFs can be readily identified using known UMFs as a target sequence or target motif with the computer-based systems described below. The presence and activity of a UMF can be confirmed by attaching the suspected UMF to a marker sequence. The resulting nucleic acid molecule is then incubated with an appropriate host under appropriate conditions and the uptake of the marker sequence is determined. As described above, a UMF will increase the frequency of uptake of a linked marker sequence.

Each of the above terms is meant to encompass all that is described for each, unless the context dictates otherwise.

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#### 4.2 NUCLEIC ACIDS OF THE INVENTION

Nucleotide sequences of the invention are set forth in the Sequence Listing.

The isolated polynucleotides of the invention include a polynucleotide comprising the nucleotide sequences of SEQ ID NO:1-1350; a polynucleotide encoding any one of the peptide sequences of SEQ ID NO:1351-2700; and a polynucleotide comprising the nucleotide sequence encoding the mature protein coding sequence of the polypeptides of any one of SEQ ID NO:1351-2700. The polynucleotides of the present invention also include, but are not limited to, a polynucleotide that hybridizes under stringent conditions to (a) the complement of any of the nucleotides sequences of SEQ ID NO:1-1350; (b) nucleotide sequences encoding any one of the amino acid sequences set forth in the Sequence Listing; (c) a polynucleotide which is an allelic variant of any polynucleotide recited above; (d) a polynucleotide which encodes a species homolog of any of the proteins recited above; or (e) a polynucleotide that encodes a polypeptide comprising a specific domain or truncation of the polypeptides of SEQ ID NO: 1351-2700. Domains of interest may depend on the nature of the encoded polypeptide; e.g., domains in receptor-like polypeptides include ligand-binding, extracellular, transmembrane, or cytoplasmic

domains, or combinations thereof; domains in immunoglobulin-like proteins include the variable immunoglobulin-like domains; domains in enzyme-like polypeptides include catalytic and substrate binding domains; and domains in ligand polypeptides include receptor-binding domains.

The polynucleotides of the invention include naturally occurring or wholly or partially synthetic DNA, *e.g.*, cDNA and genomic DNA, and RNA, *e.g.*, mRNA. The polynucleotides may include all of the coding region of the cDNA or may represent a portion of the coding region of the cDNA.

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The present invention also provides genes corresponding to the cDNA sequences disclosed herein. The corresponding genes can be isolated in accordance with known methods using the sequence information disclosed herein. Such methods include the preparation of probes or primers from the disclosed sequence information for identification and/or amplification of genes in appropriate genomic libraries or other sources of genomic materials. Further 5' and 3' sequence can be obtained using methods known in the art. For example, full length cDNA or genomic DNA that corresponds to any of the polynucleotides of SEQ ID NO:1-1350 can be obtained by screening appropriate cDNA or genomic DNA libraries under suitable hybridization conditions using any of the polynucleotides of SEQ ID NO:1-1350 or a portion thereof as a probe. Alternatively, the polynucleotides of SEQ ID NO:1-1350 may be used as the basis for suitable primer(s) that allow identification and/or amplification of genes in appropriate genomic DNA or cDNA libraries.

The nucleic acid sequences of the invention can be assembled from ESTs and sequences (including cDNA and genomic sequences) obtained from one or more public databases, such as dbEST, gbpri, and UniGene. The EST sequences can provide identifying sequence information, representative fragment or segment information, or novel segment information for the full-length gene.

The polynucleotides of the invention also provide polynucleotides including nucleotide sequences that are substantially equivalent to the polynucleotides recited above. Polynucleotides according to the invention can have, e.g., at least about 65%, at least about 70%, at least about 75%, at least about 80%, 81%, 82%, 83%, 84%, more typically at least about 85%, 86%, 87%, 88%, 89%, more typically at least about 90%, 91%, 92%, 93%, 94%, and even more typically at least about 95%, 96%, 97%, 98%, 99%, sequence identity to a polynucleotide recited above.

Included within the scope of the nucleic acid sequences of the invention are nucleic acid sequence fragments that hybridize under stringent conditions to any of the nucleotide sequences of SEQ ID NO:1-1350, or complements thereof, which fragment is greater than about 5 nucleotides, preferably 7 nucleotides, more preferably greater than 9 nucleotides and most preferably greater than 17 nucleotides. Fragments of, *e.g.* 15, 17, or 20 nucleotides or more that

are selective for (*i.e.* specifically hybridize to any one of the polynucleotides of the invention) are contemplated. Probes capable of specifically hybridizing to a polynucleotide can differentiate polynucleotide sequences of the invention from other polynucleotide sequences in the same family of genes or can differentiate human genes from genes of other species, and are preferably based on unique nucleotide sequences.

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The sequences falling within the scope of the present invention are not limited to these specific sequences, but also include allelic and species variations thereof. Allelic and species variations can be routinely determined by comparing the sequence provided SEQ ID NO:1-1350, a representative fragment thereof, or a nucleotide sequence at least 90% identical, preferably 95% identical, to SEQ ID NO:1-1350 with a sequence from another isolate of the same species. Furthermore, to accommodate codon variability, the invention includes nucleic acid molecules coding for the same amino acid sequences as do the specific ORFs disclosed herein. In other words, in the coding region of an ORF, substitution of one codon for another codon that encodes the same amino acid is expressly contemplated.

The nearest neighbor or homology result for the nucleic acids of the present invention, including SEQ ID NO:1-1350, can be obtained by searching a database using an algorithm or a program. Preferably, a BLAST which stands for Basic Local Alignment Search Tool is used to search for local sequence alignments (Altshul, S.F. J Mol. Evol. 36 290-300 (1993) and Altschul S.F. et al. J. Mol. Biol. 21:403-410 (1990)). Alternatively a FASTA version 3 search against Genpept, using Fastxy algorithm.

Species homologs (or orthologs) of the disclosed polynucleotides and proteins are also provided by the present invention. Species homologs may be isolated and identified by making suitable probes or primers from the sequences provided herein and screening a suitable nucleic acid source from the desired species.

The invention also encompasses allelic variants of the disclosed polynucleotides or proteins; that is, naturally-occurring alternative forms of the isolated polynucleotide which also encode proteins which are identical, homologous or related to that encoded by the polynucleotides.

The nucleic acid sequences of the invention are further directed to sequences which encode variants of the described nucleic acids. These amino acid sequence variants may be prepared by methods known in the art by introducing appropriate nucleotide changes into a native or variant polynucleotide. There are two variables in the construction of amino acid sequence variants: the location of the mutation and the nature of the mutation. Nucleic acids encoding the amino acid sequence variants are preferably constructed by mutating the polynucleotide to encode an amino acid sequence that does not occur in nature. These nucleic

acid alterations can be made at sites that differ in the nucleic acids from different species (variable positions) or in highly conserved regions (constant regions). Sites at such locations will typically be modified in series, *e.g.*, by substituting first with conservative choices (*e.g.*, hydrophobic amino acid to a different hydrophobic amino acid) and then with more distant choices (*e.g.*, hydrophobic amino acid to a charged amino acid), and then deletions or insertions may be made at the target site. Amino acid sequence deletions generally range from about 1 to 30 residues, preferably about 1 to 10 residues, and are typically contiguous. Amino acid insertions include amino- and/or carboxyl-terminal fusions ranging in length from one to one hundred or more residues, as well as intrasequence insertions of single or multiple amino acid residues. Intrasequence insertions may range generally from about 1 to 10 amino residues, preferably from 1 to 5 residues. Examples of terminal insertions include the heterologous signal sequences necessary for secretion or for intracellular targeting in different host cells and sequences such as FLAG or poly-histidine sequences useful for purifying the expressed protein.

In a preferred method, polynucleotides encoding the novel amino acid sequences are changed via site-directed mutagenesis. This method uses oligonucleotide sequences to alter a polynucleotide to encode the desired amino acid variant, as well as sufficient adjacent nucleotides on both sides of the changed amino acid to form a stable duplex on either side of the site of being changed. In general, the techniques of site-directed mutagenesis are well known to those of skill in the art and this technique is exemplified by publications such as, Edelman et al., DNA 2:183 (1983). A versatile and efficient method for producing site-specific changes in a polynucleotide sequence was published by Zoller and Smith, Nucleic Acids Res. 10:6487-6500 (1982). PCR may also be used to create amino acid sequence variants of the novel nucleic acids. When small amounts of template DNA are used as starting material, primer(s) that differs slightly in sequence from the corresponding region in the template DNA can generate the desired amino acid variant. PCR amplification results in a population of product DNA fragments that differ from the polynucleotide template encoding the polypeptide at the position specified by the primer. The product DNA fragments replace the corresponding region in the plasmid and this gives a polynucleotide encoding the desired amino acid variant.

A further technique for generating amino acid variants is the cassette mutagenesis technique described in Wells et al., *Gene* 34:315 (1985); and other mutagenesis techniques well known in the art, such as, for example, the techniques in Sambrook et al., supra, and *Current Protocols in Molecular Biology*, Ausubel et al. Due to the inherent degeneracy of the genetic code, other DNA sequences which encode substantially the same or a functionally equivalent amino acid sequence may be used in the practice of the invention for the cloning and expression

of these novel nucleic acids. Such DNA sequences include those which are capable of hybridizing to the appropriate novel nucleic acid sequence under stringent conditions.

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Polynucleotides encoding preferred polypeptide truncations of the invention can be used to generate polynucleotides encoding chimeric or fusion proteins comprising one or more domains of the invention and heterologous protein sequences.

The polynucleotides of the invention additionally include the complement of any of the polynucleotides recited above. The polynucleotide can be DNA (genomic, cDNA, amplified, or synthetic) or RNA. Methods and algorithms for obtaining such polynucleotides are well known to those of skill in the art and can include, for example, methods for determining hybridization conditions that can routinely isolate polynucleotides of the desired sequence identities.

In accordance with the invention, polynucleotide sequences comprising the mature protein coding sequences corresponding to any one of SEQ ID NO:1-1350, or functional equivalents thereof, may be used to generate recombinant DNA molecules that direct the expression of that nucleic acid, or a functional equivalent thereof, in appropriate host cells. Also included are the cDNA inserts of any of the clones identified herein.

A polynucleotide according to the invention can be joined to any of a variety of other nucleotide sequences by well-established recombinant DNA techniques (see Sambrook J et al. (1989) Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, NY). Useful nucleotide sequences for joining to polynucleotides include an assortment of vectors, e.g., plasmids, cosmids, lambda phage derivatives, phagemids, and the like, that are well known in the art. Accordingly, the invention also provides a vector including a polynucleotide of the invention and a host cell containing the polynucleotide. In general, the vector contains an origin of replication functional in at least one organism, convenient restriction endonuclease sites, and a selectable marker for the host cell. Vectors according to the invention include expression vectors, replication vectors, probe generation vectors, and sequencing vectors. A host cell according to the invention can be a prokaryotic or eukaryotic cell and can be a unicellular organism or part of a multicellular organism.

The present invention further provides recombinant constructs comprising a nucleic acid having any of the nucleotide sequences of SEQ ID NO:1-1350 or a fragment thereof or any other polynucleotides of the invention. In one embodiment, the recombinant constructs of the present invention comprise a vector, such as a plasmid or viral vector, into which a nucleic acid having any of the nucleotide sequences of SEQ ID NO:1-1350 or a fragment thereof is inserted, in a forward or reverse orientation. In the case of a vector comprising one of the ORFs of the present invention, the vector may further comprise regulatory sequences, including for example, a promoter, operably linked to the ORF. Large numbers of suitable vectors and promoters are

known to those of skill in the art and are commercially available for generating the recombinant constructs of the present invention. The following vectors are provided by way of example. Bacterial: pBs, phagescript, PsiX174, pBluescript SK, pBs KS, pNH8a, pNH16a, pNH18a, pNH46a (Stratagene); pTrc99A, pKK223-3, pKK233-3, pDR540, pRIT5 (Pharmacia). Eukaryotic: pWLneo, pSV2cat, pOG44, PXTI, pSG (Stratagene) pSVK3, pBPV, pMSG, pSVL (Pharmacia).

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The isolated polynucleotide of the invention may be operably linked to an expression control sequence such as the pMT2 or pED expression vectors disclosed in Kaufman et al., *Nucleic Acids Res.* 19, 4485-4490 (1991), in order to produce the protein recombinantly. Many suitable expression control sequences are known in the art. General methods of expressing recombinant proteins are also known and are exemplified in R. Kaufman, *Methods in Enzymology* 185, 537-566 (1990). As defined herein "operably linked" means that the isolated polynucleotide of the invention and an expression control sequence are situated within a vector or cell in such a way that the protein is expressed by a host cell which has been transformed (transfected) with the ligated polynucleotide/expression control sequence.

Promoter regions can be selected from any desired gene using CAT (chloramphenicol transferase) vectors or other vectors with selectable markers. Two appropriate vectors are pKK232-8 and pCM7. Particular named bacterial promoters include lacI, lacZ, T3, T7, gpt, lambda PR, and trc. Eukaryotic promoters include CMV immediate early, HSV thymidine kinase, early and late SV40, LTRs from retrovirus, and mouse metallothionein-I. Selection of the appropriate vector and promoter is well within the level of ordinary skill in the art. Generally, recombinant expression vectors will include origins of replication and selectable markers permitting transformation of the host cell, e.g., the ampicillin resistance gene of E. coli and S. cerevisiae TRP1 gene, and a promoter derived from a highly-expressed gene to direct transcription of a downstream structural sequence. Such promoters can be derived from operons encoding glycolytic enzymes such as 3-phosphoglycerate kinase (PGK), a-factor, acid phosphatase, or heat shock proteins, among others. The heterologous structural sequence is assembled in appropriate phase with translation initiation and termination sequences, and preferably, a leader sequence capable of directing secretion of translated protein into the periplasmic space or extracellular medium. Optionally, the heterologous sequence can encode a fusion protein including an amino terminal identification peptide imparting desired characteristics, e.g., stabilization or simplified purification of expressed recombinant product. Useful expression vectors for bacterial use are constructed by inserting a structural DNA sequence encoding a desired protein together with suitable translation initiation and termination signals in operable reading phase with a functional promoter. The vector will comprise one or

more phenotypic selectable markers and an origin of replication to ensure maintenance of the vector and to, if desirable, provide amplification within the host. Suitable prokaryotic hosts for transformation include *E. coli*, *Bacillus subtilis*, *Salmonella typhimurium* and various species within the genera *Pseudomonas*, *Streptomyces*, and *Staphylococcus*, although others may also be employed as a matter of choice.

As a representative but non-limiting example, useful expression vectors for bacterial use can comprise a selectable marker and bacterial origin of replication derived from commercially available plasmids comprising genetic elements of the well known cloning vector pBR322 (ATCC 37017). Such commercial vectors include, for example, pKK223-3 (Pharmacia Fine Chemicals, Uppsala, Sweden) and GEM 1 (Promega Biotech, Madison, WI, USA). These pBR322 "backbone" sections are combined with an appropriate promoter and the structural sequence to be expressed. Following transformation of a suitable host strain and growth of the host strain to an appropriate cell density, the selected promoter is induced or derepressed by appropriate means (e.g., temperature shift or chemical induction) and cells are cultured for an additional period. Cells are typically harvested by centrifugation, disrupted by physical or chemical means, and the resulting crude extract retained for further purification.

Polynucleotides of the invention can also be used to induce immune responses. For example, as described in Fan et al., *Nat. Biotech.* 17:870-872 (1999), incorporated herein by reference, nucleic acid sequences encoding a polypeptide may be used to generate antibodies against the encoded polypeptide following topical administration of naked plasmid DNA or following injection, and preferably intramuscular injection of the DNA. The nucleic acid sequences are preferably inserted in a recombinant expression vector and may be in the form of naked DNA.

### 25 **4.3 ANTISENSE**

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Another aspect of the invention pertains to isolated antisense nucleic acid molecules that are hybridizable to or complementary to the nucleic acid molecule comprising the nucleotide sequence of SEQ ID NO:1-1350, or fragments, analogs or derivatives thereof. An "antisense" nucleic acid comprises a nucleotide sequence that is complementary to a "sense" nucleic acid encoding a protein, *e.g.*, complementary to the coding strand of a double-stranded cDNA molecule or complementary to an mRNA sequence. In specific aspects, antisense nucleic acid molecules are provided that comprise a sequence complementary to at least about 10, 25, 50, 100, 250 or 500 nucleotides or an entire coding strand, or to only a portion thereof. Nucleic acid molecules encoding fragments, homologs, derivatives and analogs of a protein of any of SEQ ID

NO:1351-2700 or antisense nucleic acids complementary to a nucleic acid sequence of SEQ ID NO:1-1350 are additionally provided.

In one embodiment, an antisense nucleic acid molecule is antisense to a "coding region" of the coding strand of a nucleotide sequence of the invention. The term "coding region" refers to the region of the nucleotide sequence comprising codons which are translated into amino acid residues. In another embodiment, the antisense nucleic acid molecule is antisense to a "noncoding region" of the coding strand of a nucleotide sequence of the invention. The term "noncoding region" refers to 5' and 3' sequences which flank the coding region that are not translated into amino acids (*i.e.*, also referred to as 5' and 3' untranslated regions).

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Given the coding strand sequences encoding a nucleic acid disclosed herein (e.g., SEQ ID NO:1-1350), antisense nucleic acids of the invention can be designed according to the rules of Watson and Crick or Hoogsteen base pairing. The antisense nucleic acid molecule can be complementary to the entire coding region of a mRNA, but more preferably is an oligonucleotide that is antisense to only a portion of the coding or noncoding region of a mRNA. For example, the antisense oligonucleotide can be complementary to the region surrounding the translation start site of a mRNA. An antisense oligonucleotide can be, for example, about 5, 10, 15, 20, 25, 30, 35, 40, 45 or 50 nucleotides in length. An antisense nucleic acid of the invention can be constructed using chemical synthesis or enzymatic ligation reactions using procedures known in the art. For example, an antisense nucleic acid (e.g., an antisense oligonucleotide) can be chemically synthesized using naturally occurring nucleotides or variously modified nucleotides designed to increase the biological stability of the molecules or to increase the physical stability of the duplex formed between the antisense and sense nucleic acids, e.g., phosphorothioate derivatives and acridine substituted nucleotides can be used.

Examples of modified nucleotides that can be used to generate the antisense nucleic acid include: 5-fluorouracil, 5-bromouracil, 5-chlorouracil, 5-iodouracil, hypoxanthine, xanthine, 4-acetylcytosine, 5-(carboxyhydroxylmethyl) uracil, 5-carboxymethylaminomethyl-2-thiouridine, 5-carboxymethylaminomethyluracil, dihydrouracil, beta-D-galactosylqueosine, inosine, N6-isopentenyladenine, 1-methylguanine, 1-methylinosine, 2,2-dimethylguanine, 2-methyladenine, 2-methylguanine, 3-methylcytosine, 5-methylcytosine, N6-adenine, 7-methylguanine, 5-methylaminomethyluracil, 5-methoxyaminomethyl-2-thiouracil, beta-D-mannosylqueosine, 5'-methoxycarboxymethyluracil, 5-methoxyuracil, 2-methylthio-N6-isopentenyladenine, uracil-5-oxyacetic acid (v), wybutoxosine, pseudouracil, queosine, 2-thiocytosine, 5-methyl-2-thiouracil, 2-thiouracil, 4-thiouracil, 5-methyluracil, uracil-5-oxyacetic acid methylester, uracil-5-oxyacetic acid (v), 5-methyl-2-thiouracil, 3-(3-amino-3-N-2-carboxypropyl) uracil, (acp3)w, and 2,6-diaminopurine. Alternatively, the

antisense nucleic acid can be produced biologically using an expression vector into which a nucleic acid has been subcloned in an antisense orientation (*i.e.*, RNA transcribed from the inserted nucleic acid will be of an antisense orientation to a target nucleic acid of interest, described further in the following subsection).

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The antisense nucleic acid molecules of the invention are typically administered to a subject or generated in situ such that they hybridize with or bind to cellular mRNA and/or genomic DNA encoding a protein according to the invention to thereby inhibit expression of the protein, e.g., by inhibiting transcription and/or translation. The hybridization can be by conventional nucleotide complementarity to form a stable duplex, or, for example, in the case of an antisense nucleic acid molecule that binds to DNA duplexes, through specific interactions in the major groove of the double helix. An example of a route of administration of antisense nucleic acid molecules of the invention includes direct injection at a tissue site. Alternatively, antisense nucleic acid molecules can be modified to target selected cells and then administered systemically. For example, for systemic administration, antisense molecules can be modified such that they specifically bind to receptors or antigens expressed on a selected cell surface, e.g., by linking the antisense nucleic acid molecules to peptides or antibodies that bind to cell surface receptors or antigens. The antisense nucleic acid molecules can also be delivered to cells using the vectors described herein. To achieve sufficient intracellular concentrations of antisense molecules, vector constructs in which the antisense nucleic acid molecule is placed under the control of a strong pol II or pol III promoter are preferred.

In yet another embodiment, the antisense nucleic acid molecule of the invention is an -a nomeric nucleic acid molecule. An -a nomeric nucleic acid molecule forms specific double-stranded hybrids with complementary RNA in which, contrary to the usual -units, the strands run parallel to each other (Gaultier et al. (1987) Nucleic Acids Res 15: 6625-6641). The antisense nucleic acid molecule can also comprise a 2'-o-methylribonucleotide (Inoue et al. (1987) Nucleic Acids Res 15: 6131-6148) or a chimeric RNA -DNA analogue (Inoue et al. (1987) FEBS Lett 215: 327-330).

# 4.4 RIBOZYMES AND PNA MOIETIES

In still another embodiment, an antisense nucleic acid of the invention is a ribozyme. Ribozymes are catalytic RNA molecules with ribonuclease activity that are capable of cleaving a single-stranded nucleic acid, such as a mRNA, to which they have a complementary region. Thus, ribozymes (e.g., hammerhead ribozymes (described in Haselhoff and Gerlach (1988) *Nature* 334:585-591)) can be used to catalytically cleave a mRNA transcripts to thereby inhibit translation of a mRNA. A ribozyme having specificity for a nucleic acid of the invention can be

designed based upon the nucleotide sequence of a DNA disclosed herein (*i.e.*, SEQ ID NO:1-1350). For example, a derivative of a Tetrahymena L-19 IVS RNA can be constructed in which the nucleotide sequence of the active site is complementary to the nucleotide sequence to be cleaved in a SECX-encoding mRNA. See, *e.g.*, Cech *et al.* U.S. Pat. No. 4,987,071; and Cech *et al.* U.S. Pat. No. 5,116,742. Alternatively, SECX mRNA can be used to select a catalytic RNA having a specific ribonuclease activity from a pool of RNA molecules. See, *e.g.*, Bartel *et al.*, (1993) *Science* 261:1411-1418.

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Alternatively, gene expression can be inhibited by targeting nucleotide sequences complementary to the regulatory region (e.g., promoter and/or enhancers) to form triple helical structures that prevent transcription of the gene in target cells. See generally, Helene. (1991)

Anticancer Drug Des. 6: 569-84; Helene. et al. (1992) Ann. N.Y. Acad. Sci. 660:27-36; and Maher (1992) Bioassays 14: 807-15.

In various embodiments, the nucleic acids of the invention can be modified at the base moiety, sugar moiety or phosphate backbone to improve, e.g., the stability, hybridization, or solubility of the molecule. For example, the deoxyribose phosphate backbone of the nucleic acids can be modified to generate peptide nucleic acids (see Hyrup et al. (1996) Bioorg Med Chem 4: 5-23). As used herein, the terms "peptide nucleic acids" or "PNAs" refer to nucleic acid mimics, e.g., DNA mimics, in which the deoxyribose phosphate backbone is replaced by a pseudopeptide backbone and only the four natural nucleobases are retained. The neutral backbone of PNAs has been shown to allow for specific hybridization to DNA and RNA under conditions of low ionic strength. The synthesis of PNA oligomers can be performed using standard solid phase peptide synthesis protocols as described in Hyrup et al. (1996) above; Perry-O'Keefe et al. (1996) PNAS 93: 14670-675.

PNAs of the invention can be used in therapeutic and diagnostic applications. For example, PNAs can be used as antisense or antigene agents for sequence-specific modulation of gene expression by, *e.g.*, inducing transcription or translation arrest or inhibiting replication. PNAs of the invention can also be used, *e.g.*, in the analysis of single base pair mutations in a gene by, *e.g.*, PNA directed PCR clamping; as artificial restriction enzymes when used in combination with other enzymes, *e.g.*, S1 nucleases (Hyrup B. (1996) above); or as probes or primers for DNA sequence and hybridization (Hyrup *et al.* (1996), above; Perry-O'Keefe (1996), above).

In another embodiment, PNAs of the invention can be modified, e.g., to enhance their stability or cellular uptake, by attaching lipophilic or other helper groups to PNA, by the formation of PNA-DNA chimeras, or by the use of liposomes or other techniques of drug delivery known in the art. For example, PNA-DNA chimeras can be generated that may

combine the advantageous properties of PNA and DNA. Such chimeras allow DNA recognition enzymes, *e.g.*, RNase H and DNA polymerases, to interact with the DNA portion while the PNA portion would provide high binding affinity and specificity. PNA-DNA chimeras can be linked using linkers of appropriate lengths selected in terms of base stacking, number of bonds between the nucleobases, and orientation (Hyrup (1996) above). The synthesis of PNA-DNA chimeras can be performed as described in Hyrup (1996) above and Finn *et al.* (1996) *Nucl Acids Res* 24: 3357-63. For example, a DNA chain can be synthesized on a solid support using standard phosphoramidite coupling chemistry, and modified nucleoside analogs, *e.g.*, 5'-(4-methoxytrityl)amino-5'-deoxy-thymidine phosphoramidite, can be used between the PNA and the 5' end of DNA (Mag *et al.* (1989) *Nucl Acid Res* 17: 5973-88). PNA monomers are then coupled in a stepwise manner to produce a chimeric molecule with a 5' PNA segment and a 3' DNA segment (Finn *et al.* (1996) above). Alternatively, chimeric molecules can be synthesized with a 5' DNA segment and a 3' PNA segment. See, Petersen *et al.* (1975) *Bioorg Med Chem Lett* 5: 1119-11124.

In other embodiments, the oligonucleotide may include other appended groups such as peptides (e.g., for targeting host cell receptors *in vivo*), or agents facilitating transport across the cell membrane (see, e.g., Letsinger *et al.*, 1989, *Proc. Natl. Acad. Sci. U.S.A.* 86:6553-6556; Lemaitre *et al.*, 1987, Proc. *Natl. Acad. Sci.* 84:648-652; PCT Publication No. W088/09810) or the blood-brain barrier (see, e.g., PCT Publication No. W089/10134). In addition, oligonucleotides can be modified with hybridization triggered cleavage agents (See, *e.g.*, Krol *et al.*, 1988, *BioTechniques* 6:958-976) or intercalating agents. (See, *e.g.*, Zon, 1988, *Pharm. Res.* 5: 539-549). To this end, the oligonucleotide may be conjugated to another molecule, *e.g.*, a peptide, a hybridization triggered cross-linking agent, a transport agent, a hybridization-triggered cleavage agent, etc.

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#### **4.5 HOSTS**

The present invention further provides host cells genetically engineered to contain the polynucleotides of the invention. For example, such host cells may contain nucleic acids of the invention introduced into the host cell using known transformation, transfection or infection methods. The present invention still further provides host cells genetically engineered to express the polynucleotides of the invention, wherein such polynucleotides are in operative association with a regulatory sequence heterologous to the host cell which drives expression of the polynucleotides in the cell.

Knowledge of nucleic acid sequences allows for modification of cells to permit, or increase, expression of endogenous polypeptide. Cells can be modified (e.g., by homologous

recombination) to provide increased polypeptide expression by replacing, in whole or in part, the naturally occurring promoter with all or part of a heterologous promoter so that the cells express the polypeptide at higher levels. The heterologous promoter is inserted in such a manner that it is operatively linked to the encoding sequences. See, for example, PCT International Publication No. WO94/12650, PCT International Publication No. WO92/20808, and PCT International Publication No. WO91/09955. It is also contemplated that, in addition to heterologous promoter DNA, amplifiable marker DNA (*e.g.*, ada, dhfr, and the multifunctional CAD gene which encodes carbamyl phosphate synthase, aspartate transcarbamylase, and dihydroorotase) and/or intron DNA may be inserted along with the heterologous promoter DNA. If linked to the coding sequence, amplification of the marker DNA by standard selection methods results in coamplification of the desired protein coding sequences in the cells.

The host cell can be a higher eukaryotic host cell, such as a mammalian cell, a lower eukaryotic host cell, such as a yeast cell, or the host cell can be a prokaryotic cell, such as a bacterial cell. Introduction of the recombinant construct into the host cell can be effected by calcium phosphate transfection, DEAE, dextran mediated transfection, or electroporation (Davis, L. et al., *Basic Methods in Molecular Biology* (1986)). The host cells containing one of the polynucleotides of the invention, can be used in conventional manners to produce the gene product encoded by the isolated fragment (in the case of an ORF) or can be used to produce a heterologous protein under the control of the EMF.

Any host/vector system can be used to express one or more of the ORFs of the present invention. These include, but are not limited to, eukaryotic hosts such as HeLa cells, Cv-1 cell, COS cells, 293 cells, and Sf9 cells, as well as prokaryotic host such as *E. coli* and *B. subtilis*. The most preferred cells are those which do not normally express the particular polypeptide or protein or which expresses the polypeptide or protein at low natural level. Mature proteins can be expressed in mammalian cells, yeast, bacteria, or other cells under the control of appropriate promoters. Cell-free translation systems can also be employed to produce such proteins using RNAs derived from the DNA constructs of the present invention. Appropriate cloning and expression vectors for use with prokaryotic and eukaryotic hosts are described by Sambrook, et al., in Molecular Cloning: A Laboratory Manual, Second Edition, Cold Spring Harbor, New York (1989), the disclosure of which is hereby incorporated by reference.

Various mammalian cell culture systems can also be employed to express recombinant protein. Examples of mammalian expression systems include the COS-7 lines of monkey kidney fibroblasts, described by Gluzman, Cell 23:175 (1981). Other cell lines capable of expressing a compatible vector are, for example, the C127, monkey COS cells, Chinese Hamster Ovary (CHO) cells, human kidney 293 cells, human epidermal A431 cells, human Colo205 cells, 3T3

cells, CV-1 cells, other transformed primate cell lines, normal diploid cells, cell strains derived from *in vitro* culture of primary tissue, primary explants, HeLa cells, mouse L cells, BHK, HL-60, U937, HaK or Jurkat cells. Mammalian expression vectors will comprise an origin of replication, a suitable promoter and also any necessary ribosome binding sites, polyadenylation site, splice donor and acceptor sites, transcriptional termination sequences, and 5' flanking nontranscribed sequences. DNA sequences derived from the SV40 viral genome, for example, SV40 origin, early promoter, enhancer, splice, and polyadenylation sites may be used to provide the required nontranscribed genetic elements. Recombinant polypeptides and proteins produced in bacterial culture are usually isolated by initial extraction from cell pellets, followed by one or more salting-out, aqueous ion exchange or size exclusion chromatography steps. Protein refolding steps can be used, as necessary, in completing configuration of the mature protein. Finally, high performance liquid chromatography (HPLC) can be employed for final purification steps. Microbial cells employed in expression of proteins can be disrupted by any convenient method, including freeze-thaw cycling, sonication, mechanical disruption, or use of cell lysing agents.

Alternatively, it may be possible to produce the protein in lower eukaryotes such as yeast or insects or in prokaryotes such as bacteria. Potentially suitable yeast strains include *Saccharomyces cerevisiae*, *Schizosaccharomyces pombe*, *Kluyveromyces* strains, *Candida*, or any yeast strain capable of expressing heterologous proteins. Potentially suitable bacterial strains include *Escherichia coli*, *Bacillus subtilis*, *Salmonella typhimurium*, or any bacterial strain capable of expressing heterologous proteins. If the protein is made in yeast or bacteria, it may be necessary to modify the protein produced therein, for example by phosphorylation or glycosylation of the appropriate sites, in order to obtain the functional protein. Such covalent attachments may be accomplished using known chemical or enzymatic methods.

In another embodiment of the present invention, cells and tissues may be engineered to express an endogenous gene comprising the polynucleotides of the invention under the control of inducible regulatory elements, in which case the regulatory sequences of the endogenous gene may be replaced by homologous recombination. As described herein, gene targeting can be used to replace a gene's existing regulatory region with a regulatory sequence isolated from a different gene or a novel regulatory sequence synthesized by genetic engineering methods. Such regulatory sequences may be comprised of promoters, enhancers, scaffold-attachment regions, negative regulatory elements, transcriptional initiation sites, regulatory protein binding sites or combinations of said sequences. Alternatively, sequences which affect the structure or stability of the RNA or protein produced may be replaced, removed, added, or otherwise modified by targeting. These sequence include polyadenylation signals, mRNA stability elements, splice

sites, leader sequences for enhancing or modifying transport or secretion properties of the protein, or other sequences which alter or improve the function or stability of protein or RNA molecules.

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The targeting event may be a simple insertion of the regulatory sequence, placing the gene under the control of the new regulatory sequence, e.g., inserting a new promoter or enhancer or both upstream of a gene. Alternatively, the targeting event may be a simple deletion of a regulatory element, such as the deletion of a tissue-specific negative regulatory element. Alternatively, the targeting event may replace an existing element; for example, a tissue-specific enhancer can be replaced by an enhancer that has broader or different cell-type specificity than the naturally occurring elements. Here, the naturally occurring sequences are deleted and new sequences are added. In all cases, the identification of the targeting event may be facilitated by the use of one or more selectable marker genes that are contiguous with the targeting DNA, allowing for the selection of cells in which the exogenous DNA has integrated into the host cell genome. The identification of the targeting event may also be facilitated by the use of one or more marker genes exhibiting the property of negative selection, such that the negatively selectable marker is linked to the exogenous DNA, but configured such that the negatively selectable marker flanks the targeting sequence, and such that a correct homologous recombination event with sequences in the host cell genome does not result in the stable integration of the negatively selectable marker. Markers useful for this purpose include the Herpes Simplex Virus thymidine kinase (TK) gene or the bacterial xanthine-guanine phosphoribosyl-transferase (gpt) gene.

The gene targeting or gene activation techniques which can be used in accordance with this aspect of the invention are more particularly described in U.S. Patent No. 5,272,071 to Chappel; U.S. Patent No. 5,578,461 to Sherwin et al.; International Application No. PCT/US92/09627 (WO93/09222) by Selden et al.; and International Application No. PCT/US90/06436 (WO91/06667) by Skoultchi et al., each of which is incorporated by reference herein in its entirety.

# 4.6 POLYPEPTIDES OF THE INVENTION

The isolated polypeptides of the invention include, but are not limited to, a polypeptide comprising: the amino acid sequences set forth as any one of SEQ ID NO:1351-2700 or an amino acid sequence encoded by any one of the nucleotide sequences SEQ ID NO:1-1350 or the corresponding full length or mature protein. Polypeptides of the invention also include polypeptides preferably with biological or immunological activity that are encoded by: (a) a polynucleotide having any one of the nucleotide sequences set forth in SEQ ID NO:1-1350 or (b)

polynucleotides encoding any one of the amino acid sequences set forth as SEQ ID NO:1351-2700 or (c) polynucleotides that hybridize to the complement of the polynucleotides of either (a) or (b) under stringent hybridization conditions. The invention also provides biologically active or immunologically active variants of any of the amino acid sequences set forth as SEQ ID NO:1351-2700 or the corresponding full length or mature protein; and "substantial equivalents" thereof (e.g., with at least about 65%, at least about 70%, at least about 75%, at least about 80%, at least about 85%, 86%, 87%, 88%, 89%, at least about 90%, 91%, 92%, 93%, 94%, typically at least about 95%, 96%, 97%, more typically at least about 98%, or most typically at least about 99% amino acid identity) that retain biological activity. Polypeptides encoded by allelic variants may have a similar, increased, or decreased activity compared to polypeptides comprising SEQ ID NO:1351-2700.

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Fragments of the proteins of the present invention which are capable of exhibiting biological activity are also encompassed by the present invention. Fragments of the protein may be in linear form or they may be cyclized using known methods, for example, as described in H. U. Saragovi, et al., Bio/Technology 10, 773-778 (1992) and in R. S. McDowell, et al., J. Amer. Chem. Soc. 114, 9245-9253 (1992), both of which are incorporated herein by reference. Such fragments may be fused to carrier molecules such as immunoglobulins for many purposes, including increasing the valency of protein binding sites.

The present invention also provides both full-length and mature forms (for example, without a signal sequence or precursor sequence) of the disclosed proteins. The protein coding sequence is identified in the sequence listing by translation of the disclosed nucleotide sequences. The mature form of such protein may be obtained by expression of a full-length polynucleotide in a suitable mammalian cell or other host cell. The sequence of the mature form of the protein is also determinable from the amino acid sequence of the full-length form. Where proteins of the present invention are membrane bound, soluble forms of the proteins are also provided. In such forms, part or all of the regions causing the proteins to be membrane bound are deleted so that the proteins are fully secreted from the cell in which they are expressed.

Protein compositions of the present invention may further comprise an acceptable carrier, such as a hydrophilic, *e.g.*, pharmaceutically acceptable, carrier.

The present invention further provides isolated polypeptides encoded by the nucleic acid fragments of the present invention or by degenerate variants of the nucleic acid fragments of the present invention. By "degenerate variant" is intended nucleotide fragments which differ from a nucleic acid fragment of the present invention (e.g., an ORF) by nucleotide sequence but, due to the degeneracy of the genetic code, encode an identical polypeptide sequence. Preferred nucleic acid fragments of the present invention are the ORFs that encode proteins.

A variety of methodologies known in the art can be utilized to obtain any one of the isolated polypeptides or proteins of the present invention. At the simplest level, the amino acid sequence can be synthesized using commercially available peptide synthesizers. The synthetically-constructed protein sequences, by virtue of sharing primary, secondary or tertiary structural and/or conformational characteristics with proteins may possess biological properties in common therewith, including protein activity. This technique is particularly useful in producing small peptides and fragments of larger polypeptides. Fragments are useful, for example, in generating antibodies against the native polypeptide. Thus, they may be employed as biologically active or immunological substitutes for natural, purified proteins in screening of therapeutic compounds and in immunological processes for the development of antibodies.

The polypeptides and proteins of the present invention can alternatively be purified from cells which have been altered to express the desired polypeptide or protein. As used herein, a cell is said to be altered to express a desired polypeptide or protein when the cell, through genetic manipulation, is made to produce a polypeptide or protein which it normally does not produce or which the cell normally produces at a lower level. One skilled in the art can readily adapt procedures for introducing and expressing either recombinant or synthetic sequences into eukaryotic or prokaryotic cells in order to generate a cell which produces one of the polypeptides or proteins of the present invention.

The invention also relates to methods for producing a polypeptide comprising growing a culture of host cells of the invention in a suitable culture medium, and purifying the protein from the cells or the culture in which the cells are grown. For example, the methods of the invention include a process for producing a polypeptide in which a host cell containing a suitable expression vector that includes a polynucleotide of the invention is cultured under conditions that allow expression of the encoded polypeptide. The polypeptide can be recovered from the culture, conveniently from the culture medium, or from a lysate prepared from the host cells and further purified. Preferred embodiments include those in which the protein produced by such process is a full length or mature form of the protein.

In an alternative method, the polypeptide or protein is purified from bacterial cells which naturally produce the polypeptide or protein. One skilled in the art can readily follow known methods for isolating polypeptides and proteins in order to obtain one of the isolated polypeptides or proteins of the present invention. These include, but are not limited to, immunochromatography, HPLC, size-exclusion chromatography, ion-exchange chromatography, and immuno-affinity chromatography. See, e.g., Scopes, Protein Purification: Principles and Practice, Springer-Verlag (1994); Sambrook, et al., in Molecular Cloning: A Laboratory Manual; Ausubel et al., Current Protocols in Molecular Biology. Polypeptide fragments that

retain biological/immunological activity include fragments comprising greater than about 100 amino acids, or greater than about 200 amino acids, and fragments that encode specific protein domains.

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The purified polypeptides can be used in *in vitro* binding assays which are well known in the art to identify molecules which bind to the polypeptides. These molecules include but are not limited to, for *e.g.*, small molecules, molecules from combinatorial libraries, antibodies or other proteins. The molecules identified in the binding assay are then tested for antagonist or agonist activity in *in vivo* tissue culture or animal models that are well known in the art. In brief, the molecules are titrated into a plurality of cell cultures or animals and then tested for either cell/animal death or prolonged survival of the animal/cells.

In addition, the peptides of the invention or molecules capable of binding to the peptides may be complexed with toxins, *e.g.*, ricin or cholera, or with other compounds that are toxic to cells. The toxin-binding molecule complex is then targeted to a tumor or other cell by the specificity of the binding molecule for SEQ ID NO:1351-2700.

The protein of the invention may also be expressed as a product of transgenic animals, e.g., as a component of the milk of transgenic cows, goats, pigs, or sheep which are characterized by somatic or germ cells containing a nucleotide sequence encoding the protein.

The proteins provided herein also include proteins characterized by amino acid sequences similar to those of purified proteins but into which modification are naturally provided or deliberately engineered. For example, modifications, in the peptide or DNA sequence, can be made by those skilled in the art using known techniques. Modifications of interest in the protein sequences may include the alteration, substitution, replacement, insertion or deletion of a selected amino acid residue in the coding sequence. For example, one or more of the cysteine residues may be deleted or replaced with another amino acid to alter the conformation of the molecule. Techniques for such alteration, substitution, replacement, insertion or deletion are well known to those skilled in the art (see, e.g., U.S. Pat. No. 4,518,584). Preferably, such alteration, substitution, replacement, insertion or deletion retains the desired activity of the protein. Regions of the protein that are important for the protein function can be determined by various methods known in the art including the alanine-scanning method which involved systematic substitution of single or strings of amino acids with alanine, followed by testing the resulting alanine-containing variant for biological activity. This type of analysis determines the importance of the substituted amino acid(s) in biological activity. Regions of the protein that are important for protein function may be determined by the eMATRIX program.

Other fragments and derivatives of the sequences of proteins which would be expected to retain protein activity in whole or in part and are useful for screening or other immunological

methodologies may also be easily made by those skilled in the art given the disclosures herein. Such modifications are encompassed by the present invention.

The protein may also be produced by operably linking the isolated polynucleotide of the invention to suitable control sequences in one or more insect expression vectors, and employing an insect expression system. Materials and methods for baculovirus/insect cell expression systems are commercially available in kit form from, *e.g.*, Invitrogen, San Diego, Calif., U.S.A. (the MaxBat<sup>TM</sup> kit), and such methods are well known in the art, as described in Summers and Smith, Texas Agricultural Experiment Station Bulletin No. 1555 (1987), incorporated herein by reference. As used herein, an insect cell capable of expressing a polynucleotide of the present invention is "transformed."

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The protein of the invention may be prepared by culturing transformed host cells under culture conditions suitable to express the recombinant protein. The resulting expressed protein may then be purified from such culture (*i.e.*, from culture medium or cell extracts) using known purification processes, such as gel filtration and ion exchange chromatography. The purification of the protein may also include an affinity column containing agents which will bind to the protein; one or more column steps over such affinity resins as concanavalin A-agarose, heparin-toyopearl<sup>TM</sup> or Cibacrom blue 3GA Sepharose<sup>TM</sup>; one or more steps involving hydrophobic interaction chromatography using such resins as phenyl ether, butyl ether, or propyl ether; or immunoaffinity chromatography.

Alternatively, the protein of the invention may also be expressed in a form which will facilitate purification. For example, it may be expressed as a fusion protein, such as those of maltose binding protein (MBP), glutathione-S-transferase (GST) or thioredoxin (TRX), or as a His tag. Kits for expression and purification of such fusion proteins are commercially available from New England BioLab (Beverly, Mass.), Pharmacia (Piscataway, N.J.) and Invitrogen, respectively. The protein can also be tagged with an epitope and subsequently purified by using a specific antibody directed to such epitope. One such epitope ("FLAG®") is commercially available from Kodak (New Haven, Conn.).

Finally, one or more reverse-phase high performance liquid chromatography (RP-HPLC) steps employing hydrophobic RP-HPLC media, e.g., silica gel having pendant methyl or other aliphatic groups, can be employed to further purify the protein. Some or all of the foregoing purification steps, in various combinations, can also be employed to provide a substantially homogeneous isolated recombinant protein. The protein thus purified is substantially free of other mammalian proteins and is defined in accordance with the present invention as an "isolated protein."

The polypeptides of the invention include analogs (variants). This embraces fragments, as well as peptides in which one or more amino acids has been deleted, inserted, or substituted. Also, analogs of the polypeptides of the invention embrace fusions of the polypeptides or modifications of the polypeptides of the invention, wherein the polypeptide or analog is fused to another moiety or moieties, *e.g.*, targeting moiety or another therapeutic agent. Such analogs may exhibit improved properties such as activity and/or stability. Examples of moieties which may be fused to the polypeptide or an analog include, for example, targeting moieties which provide for the delivery of polypeptide to pancreatic cells, *e.g.*, antibodies to pancreatic cells, antibodies to immune cells such as T-cells, monocytes, dendritic cells, granulocytes, etc., as well as receptor and ligands expressed on pancreatic or immune cells. Other moieties which may be fused to the polypeptide include therapeutic agents which are used for treatment, for example, immunosuppressive drugs such as cyclosporin, SK506, azathioprine, CD3 antibodies and steroids. Also, polypeptides may be fused to immune modulators, and other cytokines such as alpha or beta interferon.

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# 4.6.1 DETERMINING POLYPEPTIDE AND POLYNUCLEOTIDE IDENTITY AND SIMILARITY

Preferred identity and/or similarity are designed to give the largest match between the sequences tested. Methods to determine identity and similarity are codified in computer 20 programs including, but are not limited to, the GCG program package, including GAP (Devereux, J., et al., Nucleic Acids Research 12(1):387 (1984); Genetics Computer Group, University of Wisconsin, Madison, WI), BLASTP, BLASTN, BLASTX, FASTA (Altschul, S.F. et al., J. Molec. Biol. 215:403-410 (1990), PSI-BLAST (Altschul S.F. et al., Nucleic Acids Res. vol. 25, pp. 3389-3402, herein incorporated by reference), eMatrix software (Wu et al., J. Comp. 25 Biol., Vol. 6, pp. 219-235 (1999), herein incorporated by reference), eMotif software (Nevill-Manning et al, ISMB-97, Vol. 4, pp. 202-209, herein incorporated by reference), pFam software (Sonnhammer et al., Nucleic Acids Res., Vol. 26(1), pp. 320-322 (1998), herein incorporated by reference) and the Kyte-Doolittle hydrophobocity prediction algorithm (J. Mol Biol, 157, pp. 105-31 (1982), incorporated herein by reference). The BLAST programs are publicly available 30 from the National Center for Biotechnology Information (NCBI) and other sources (BLAST Manual, Altschul, S., et al. NCB NLM NIH Bethesda, MD 20894; Altschul, S., et al., J. Mol. Biol. 215:403-410 (1990).

# 4.7 CHIMERIC AND FUSION PROTEINS

The invention also provides chimeric or fusion proteins. As used herein, a "chimeric protein" or "fusion protein" comprises a polypeptide of the invention operatively linked to

another polypeptide. Within a fusion protein the polypeptide according to the invention can correspond to all or a portion of a protein according to the invention. In one embodiment, a fusion protein comprises at least one biologically active portion of a protein according to the invention. In another embodiment, a fusion protein comprises at least two biologically active portions of a protein according to the invention. Within the fusion protein, the term "operatively linked" is intended to indicate that the polypeptide according to the invention and the other polypeptide are fused in-frame to each other. The polypeptide can be fused to the N-terminus or C-terminus.

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For example, in one embodiment a fusion protein comprises a polypeptide according to the invention operably linked to the extracellular domain of a second protein.

In another embodiment, the fusion protein is a GST-fusion protein in which the polypeptide sequences of the invention are fused to the C-terminus of the GST (*i.e.*, glutathione S-transferase) sequences.

In another embodiment, the fusion protein is an immunoglobulin fusion protein in which the polypeptide sequences according to the invention comprises one or more domains are fused to sequences derived from a member of the immunoglobulin protein family. The immunoglobulin fusion proteins of the invention can be incorporated into pharmaceutical compositions and administered to a subject to inhibit an interaction between a ligand and a protein of the invention on the surface of a cell, to thereby suppress signal transduction *in vivo*. The immunoglobulin fusion proteins can be used to affect the bioavailability of a cognate ligand. Inhibition of the ligand/protein interaction may be useful therapeutically for both the treatment of proliferative and differentiative disorders, *e,g.*, cancer as well as modulating (*e.g.*, promoting or inhibiting) cell survival. Moreover, the immunoglobulin fusion proteins of the invention can be used as immunogens to produce antibodies in a subject, to purify ligands, and in screening assays to identify molecules that inhibit the interaction of a polypeptide of the invention with a ligand.

A chimeric or fusion protein of the invention can be produced by standard recombinant DNA techniques. For example, DNA fragments coding for the different polypeptide sequences are ligated together in-frame in accordance with conventional techniques, *e.g.*, by employing blunt-ended or stagger-ended termini for ligation, restriction enzyme digestion to provide for appropriate termini, filling-in of cohesive ends as appropriate, alkaline phosphatase treatment to avoid undesirable joining, and enzymatic ligation. In another embodiment, the fusion gene can be synthesized by conventional techniques including automated DNA synthesizers. Alternatively, PCR amplification of gene fragments can be carried out using anchor primers that give rise to complementary overhangs between two consecutive gene fragments that can subsequently be annealed and reamplified to generate a chimeric gene sequence (see, for

example, Ausubel et al. (eds.) Current Protocols in Molecular Biology, John Wiley & Sons, 1992). Moreover, many expression vectors are commercially available that already encode a fusion moiety (e.g., a GST polypeptide). A nucleic acid encoding a polypeptide of the invention can be cloned into such an expression vector such that the fusion moiety is linked in-frame to the protein of the invention.

# 4.8 GENE THERAPY

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Mutations in the polynucleotides of the invention gene may result in loss of normal function of the encoded protein. The invention thus provides gene therapy to restore normal activity of the polypeptides of the invention; or to treat disease states involving polypeptides of the invention. Delivery of a functional gene encoding polypeptides of the invention to appropriate cells is effected ex vivo, in situ, or in vivo by use of vectors, and more particularly viral vectors (e.g., adenovirus, adeno-associated virus, or a retrovirus), or ex vivo by use of physical DNA transfer methods (e.g., liposomes or chemical treatments). See, for example, Anderson, Nature, supplement to vol. 392, no. 6679, pp.25-20 (1998). For additional reviews of gene therapy technology see Friedmann, Science, 244: 1275-1281 (1989); Verma, Scientific American: 68-84 (1990); and Miller, Nature, 357: 455-460 (1992). Introduction of any one of the nucleotides of the present invention or a gene encoding the polypeptides of the present invention can also be accomplished with extrachromosomal substrates (transient expression) or artificial chromosomes (stable expression). Cells may also be cultured ex vivo in the presence of proteins of the present invention in order to proliferate or to produce a desired effect on or activity in such cells. Treated cells can then be introduced in vivo for therapeutic purposes. Alternatively, it is contemplated that in other human disease states, preventing the expression of or inhibiting the activity of polypeptides of the invention will be useful in treating the disease states. It is contemplated that antisense therapy or gene therapy could be applied to negatively regulate the expression of polypeptides of the invention.

Other methods inhibiting expression of a protein include the introduction of antisense molecules to the nucleic acids of the present invention, their complements, or their translated RNA sequences, by methods known in the art. Further, the polypeptides of the present invention can be inhibited by using targeted deletion methods, or the insertion of a negative regulatory element such as a silencer, which is tissue specific.

The present invention still further provides cells genetically engineered *in vivo* to express the polynucleotides of the invention, wherein such polynucleotides are in operative association with a regulatory sequence heterologous to the host cell which drives expression of the polynucleotides in

the cell. These methods can be used to increase or decrease the expression of the polynucleotides of the present invention.

Knowledge of DNA sequences provided by the invention allows for modification of cells to permit, increase, or decrease, expression of endogenous polypeptide. Cells can be modified (e.g., by homologous recombination) to provide increased polypeptide expression by replacing, in whole or in part, the naturally occurring promoter with all or part of a heterologous promoter so that the cells express the protein at higher levels. The heterologous promoter is inserted in such a manner that it is operatively linked to the desired protein encoding sequences. See, for example, PCT International Publication No. WO 94/12650, PCT International Publication No. WO 92/20808, and PCT International Publication No. WO 91/09955. It is also contemplated that, in addition to heterologous promoter DNA, amplifiable marker DNA (e.g., ada, dhfr, and the multifunctional CAD gene which encodes carbamyl phosphate synthase, aspartate transcarbamylase, and dihydroorotase) and/or intron DNA may be inserted along with the heterologous promoter DNA. If linked to the desired protein coding sequence, amplification of the marker DNA by standard selection methods results in co-amplification of the desired protein coding sequences in the cells.

In another embodiment of the present invention, cells and tissues may be engineered to express an endogenous gene comprising the polynucleotides of the invention under the control of inducible regulatory elements, in which case the regulatory sequences of the endogenous gene may be replaced by homologous recombination. As described herein, gene targeting can be used to replace a gene's existing regulatory region with a regulatory sequence isolated from a different gene or a novel regulatory sequence synthesized by genetic engineering methods. Such regulatory sequences may be comprised of promoters, enhancers, scaffold-attachment regions, negative regulatory elements, transcriptional initiation sites, regulatory protein binding sites or combinations of said sequences. Alternatively, sequences which affect the structure or stability of the RNA or protein produced may be replaced, removed, added, or otherwise modified by targeting. These sequences include polyadenylation signals, mRNA stability elements, splice sites, leader sequences for enhancing or modifying transport or secretion properties of the protein, or other sequences which alter or improve the function or stability of protein or RNA molecules.

The targeting event may be a simple insertion of the regulatory sequence, placing the gene under the control of the new regulatory sequence, e.g., inserting a new promoter or enhancer or both upstream of a gene. Alternatively, the targeting event may be a simple deletion of a regulatory element, such as the deletion of a tissue-specific negative regulatory element. Alternatively, the targeting event may replace an existing element; for example, a tissue-specific enhancer can be replaced by an enhancer that has broader or different cell-type specificity than the naturally occurring elements. Here, the naturally occurring sequences are deleted and new sequences are

added. In all cases, the identification of the targeting event may be facilitated by the use of one or more selectable marker genes that are contiguous with the targeting DNA, allowing for the selection of cells in which the exogenous DNA has integrated into the cell genome. The identification of the targeting event may also be facilitated by the use of one or more marker genes exhibiting the property of negative selection, such that the negatively selectable marker is linked to the exogenous DNA, but configured such that the negatively selectable marker flanks the targeting sequence, and such that a correct homologous recombination event with sequences in the host cell genome does not result in the stable integration of the negatively selectable marker. Markers useful for this purpose include the Herpes Simplex Virus thymidine kinase (TK) gene or the bacterial xanthine-guanine phosphoribosyl-transferase (gpt) gene.

The gene targeting or gene activation techniques which can be used in accordance with this aspect of the invention are more particularly described in U.S. Patent No. 5,272,071 to Chappel; U.S. Patent No. 5,578,461 to Sherwin et al.; International Application No. PCT/US92/09627 (WO93/09222) by Selden et al.; and International Application No. PCT/US90/06436 (WO91/06667) by Skoultchi et al., each of which is incorporated by reference herein in its entirety.

# 4.9 TRANSGENIC ANIMALS

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In preferred methods to determine biological functions of the polypeptides of the invention in vivo, one or more genes provided by the invention are either over expressed or inactivated in the germ line of animals using homologous recombination [Capecchi, Science 244:1288-1292 (1989)]. Animals in which the gene is over expressed, under the regulatory control of exogenous or endogenous promoter elements, are known as transgenic animals. Animals in which an endogenous gene has been inactivated by homologous recombination are referred to as "knockout" animals. Knockout animals, preferably non-human mammals, can be prepared as described in U.S. Patent No. 5,557,032, incorporated herein by reference. Transgenic animals are useful to determine the roles polypeptides of the invention play in biological processes, and preferably in disease states. Transgenic animals are useful as model systems to identify compounds that modulate lipid metabolism. Transgenic animals, preferably non-human mammals, are produced using methods as described in U.S. Patent No 5,489,743 and PCT Publication No. WO94/28122, incorporated herein by reference.

Transgenic animals can be prepared wherein all or part of a promoter of the polynucleotides of the invention is either activated or inactivated to alter the level of expression of the polypeptides of the invention. Inactivation can be carried out using homologous recombination methods described above. Activation can be achieved by supplementing or even replacing the homologous promoter to provide for increased protein expression. The homologous

promoter can be supplemented by insertion of one or more heterologous enhancer elements known to confer promoter activation in a particular tissue.

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The polynucleotides of the present invention also make possible the development, through, *e.g.*, homologous recombination or knock out strategies, of animals that fail to express polypeptides of the invention or that express a variant polypeptide. Such animals are useful as models for studying the *in vivo* activities of polypeptide as well as for studying modulators of the polypeptides of the invention.

In preferred methods to determine biological functions of the polypeptides of the invention *in vivo*, one or more genes provided by the invention are either over expressed or inactivated in the germ line of animals using homologous recombination [Capecchi, Science 244:1288-1292 (1989)]. Animals in which the gene is over expressed, under the regulatory control of exogenous or endogenous promoter elements, are known as transgenic animals. Animals in which an endogenous gene has been inactivated by homologous recombination are referred to as "knockout" animals. Knockout animals, preferably non-human mammals, can be prepared as described in U.S. Patent No. 5,557,032, incorporated herein by reference. Transgenic animals are useful to determine the roles polypeptides of the invention play in biological processes, and preferably in disease states. Transgenic animals are useful as model systems to identify compounds that modulate lipid metabolism. Transgenic animals, preferably non-human mammals, are produced using methods as described in U.S. Patent No 5,489,743 and PCT Publication No. WO94/28122, incorporated herein by reference.

Transgenic animals can be prepared wherein all or part of the polynucleotides of the invention promoter is either activated or inactivated to alter the level of expression of the polypeptides of the invention. Inactivation can be carried out using homologous recombination methods described above. Activation can be achieved by supplementing or even replacing the homologous promoter to provide for increased protein expression. The homologous promoter can be supplemented by insertion of one or more heterologous enhancer elements known to confer promoter activation in a particular tissue.

### 4.10 USES AND BIOLOGICAL ACTIVITY

The polynucleotides and proteins of the present invention are expected to exhibit one or more of the uses or biological activities (including those associated with assays cited herein) identified herein. Uses or activities described for proteins of the present invention may be provided by administration or use of such proteins or of polynucleotides encoding such proteins (such as, for example, in gene therapies or vectors suitable for introduction of DNA). The mechanism underlying the particular condition or pathology will dictate whether the

polypeptides of the invention, the polynucleotides of the invention or modulators (activators or inhibitors) thereof would be beneficial to the subject in need of treatment. Thus, "therapeutic compositions of the invention" include compositions comprising isolated polynucleotides (including recombinant DNA molecules, cloned genes and degenerate variants thereof) or polypeptides of the invention (including full length protein, mature protein and truncations or domains thereof), or compounds and other substances that modulate the overall activity of the target gene products, either at the level of target gene/protein expression or target protein activity. Such modulators include polypeptides, analogs, (variants), including fragments and fusion proteins, antibodies and other binding proteins; chemical compounds that directly or indirectly activate or inhibit the polypeptides of the invention (identified, *e.g.*, via drug screening assays as described herein); antisense polynucleotides and polynucleotides suitable for triple helix formation; and in particular antibodies or other binding partners that specifically recognize one or more epitopes of the polypeptides of the invention.

The polypeptides of the present invention may likewise be involved in cellular activation or in one of the other physiological pathways described herein.

## 4.10.1 RESEARCH USES AND UTILITIES

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The polynucleotides provided by the present invention can be used by the research community for various purposes. The polynucleotides can be used to express recombinant protein for analysis, characterization or therapeutic use; as markers for tissues in which the corresponding protein is preferentially expressed (either constitutively or at a particular stage of tissue differentiation or development or in disease states); as molecular weight markers on gels; as chromosome markers or tags (when labeled) to identify chromosomes or to map related gene positions; to compare with endogenous DNA sequences in patients to identify potential genetic disorders; as probes to hybridize and thus discover novel, related DNA sequences; as a source of information to derive PCR primers for genetic fingerprinting; as a probe to "subtract-out" known sequences in the process of discovering other novel polynucleotides; for selecting and making oligomers for attachment to a "gene chip" or other support, including for examination of expression patterns; to raise anti-protein antibodies using DNA immunization techniques; and as an antigen to raise anti-DNA antibodies or elicit another immune response. Where the polynucleotide encodes a protein which binds or potentially binds to another protein (such as, for example, in a receptor-ligand interaction), the polynucleotide can also be used in interaction trap assays (such as, for example, that described in Gyuris et al., Cell 75:791-803 (1993)) to identify polynucleotides encoding the other protein with which binding occurs or to identify inhibitors of the binding interaction.

The polypeptides provided by the present invention can similarly be used in assays to determine biological activity, including in a panel of multiple proteins for high-throughput screening; to raise antibodies or to elicit another immune response; as a reagent (including the labeled reagent) in assays designed to quantitatively determine levels of the protein (or its receptor) in biological fluids; as markers for tissues in which the corresponding polypeptide is preferentially expressed (either constitutively or at a particular stage of tissue differentiation or development or in a disease state); and, of course, to isolate correlative receptors or ligands. Proteins involved in these binding interactions can also be used to screen for peptide or small molecule inhibitors or agonists of the binding interaction.

Any or all of these research utilities are capable of being developed into reagent grade or kit format for commercialization as research products.

Methods for performing the uses listed above are well known to those skilled in the art. References disclosing such methods include without limitation "Molecular Cloning: A Laboratory Manual", 2d ed., Cold Spring Harbor Laboratory Press, Sambrook, J., E. F. Fritsch and T. Maniatis eds., 1989, and "Methods in Enzymology: Guide to Molecular Cloning Techniques", Academic Press, Berger, S. L. and A. R. Kimmel eds., 1987.

## 4.10.2 NUTRITIONAL USES

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Polynucleotides and polypeptides of the present invention can also be used as nutritional sources or supplements. Such uses include without limitation use as a protein or amino acid supplement, use as a carbon source, use as a nitrogen source and use as a source of carbohydrate. In such cases the polypeptide or polynucleotide of the invention can be added to the feed of a particular organism or can be administered as a separate solid or liquid preparation, such as in the form of powder, pills, solutions, suspensions or capsules. In the case of microorganisms, the polypeptide or polynucleotide of the invention can be added to the medium in or on which the microorganism is cultured.

# 4.10.3 CYTOKINE AND CELL PROLIFERATION/DIFFERENTIATION ACTIVITY

A polypeptide of the present invention may exhibit activity relating to cytokine, cell proliferation (either inducing or inhibiting) or cell differentiation (either inducing or inhibiting) activity or may induce production of other cytokines in certain cell populations. A polynucleotide of the invention can encode a polypeptide exhibiting such attributes. Many protein factors discovered to date, including all known cytokines, have exhibited activity in one or more factor-dependent cell proliferation assays, and hence the assays serve as a convenient

confirmation of cytokine activity. The activity of therapeutic compositions of the present invention is evidenced by any one of a number of routine factor dependent cell proliferation assays for cell lines including, without limitation, 32D, DA2, DA1G, T10, B9, B9/11, BaF3, MC9/G, M+(preB M+), 2E8, RB5, DA1, 123, T1165, HT2, CTLL2, TF-1, Mo7e, CMK, HUVEC, and Caco. Therapeutic compositions of the invention can be used in the following:

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Assays for T-cell or thymocyte proliferation include without limitation those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, *In Vitro* assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Takai et al., J. Immunol. 137:3494-3500, 1986; Bertagnolli et al., J. Immunol. 145:1706-1712, 1990; Bertagnolli et al., Cellular Immunology 133:327-341, 1991; Bertagnolli, et al., I. Immunol. 149:3778-3783, 1992; Bowman et al., I. Immunol. 152:1756-1761, 1994.

Assays for cytokine production and/or proliferation of spleen cells, lymph node cells or thymocytes include, without limitation, those described in: Polyclonal T cell stimulation, Kruisbeek, A. M. and Shevach, E. M. In Current Protocols in Immunology. J. E. e.a. Coligan eds. Vol 1 pp. 3.12.1-3.12.14, John Wiley and Sons, Toronto. 1994; and Measurement of mouse and human interleukin-γ, Schreiber, R. D. In Current Protocols in Immunology. J. E. e.a. Coligan eds. Vol 1 pp. 6.8.1-6.8.8, John Wiley and Sons, Toronto. 1994.

Assays for proliferation and differentiation of hematopoietic and lymphopoietic cells include, without limitation, those described in: Measurement of Human and Murine Interleukin 2 and Interleukin 4, Bottomly, K., Davis, L. S. and Lipsky, P. E. In Current Protocols in Immunology. J. E. e.a. Coligan eds. Vol 1 pp. 6.3.1-6.3.12, John Wiley and Sons, Toronto. 1991; deVries et al., J. Exp. Med. 173:1205-1211, 1991; Moreau et al., Nature 336:690-692, 1988; Greenberger et al., Proc. Natl. Acad. Sci. U.S.A. 80:2931-2938, 1983; Measurement of mouse and human interleukin 6--Nordan, R. In Current Protocols in Immunology. J. E. Coligan eds. Vol 1 pp. 6.6.1-6.6.5, John Wiley and Sons, Toronto. 1991; Smith et al., Proc. Natl. Aced. Sci. U.S.A. 83:1857-1861, 1986; Measurement of human Interleukin 11--Bennett, F., Giannotti, J., Clark, S. C. and Turner, K. J. In Current Protocols in Immunology. J. E. Coligan eds. Vol 1 pp. 6.15.1 John Wiley and Sons, Toronto. 1991; Measurement of mouse and human Interleukin 9--Ciarletta, A., Giannotti, J., Clark, S. C. and Turner, K. J. In Current Protocols in Immunology. J. E. Coligan eds. Vol 1 pp. 6.13.1, John Wiley and Sons, Toronto. 1991.

Assays for T-cell clone responses to antigens (which will identify, among others, proteins that affect APC-T cell interactions as well as direct T-cell effects by measuring proliferation and cytokine production) include, without limitation, those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W Strober,

Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, *In Vitro* assays for Mouse Lymphocyte Function; Chapter 6, Cytokines and their cellular receptors; Chapter 7, Immunologic studies in Humans); Weinberger et al., Proc. Natl. Acad. Sci. USA 77:6091-6095, 1980; Weinberger et al., Eur. J. Immun. 11:405-411, 1981; Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988.

## 4.10.4 STEM CELL GROWTH FACTOR ACTIVITY

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A polypeptide of the present invention may exhibit stem cell growth factor activity and be involved in the proliferation, differentiation and survival of pluripotent and totipotent stem cells including primordial germ cells, embryonic stem cells, hematopoietic stem cells and/or germ line stem cells. Administration of the polypeptide of the invention to stem cells *in vivo* or *ex vivo* is expected to maintain and expand cell populations in a totipotential or pluripotential state which would be useful for re-engineering damaged or diseased tissues, transplantation, manufacture of bio-pharmaceuticals and the development of bio-sensors. The ability to produce large quantities of human cells has important working applications for the production of human proteins which currently must be obtained from non-human sources or donors, implantation of cells to treat diseases such as Parkinson's, Alzheimer's and other neurodegenerative diseases; tissues for grafting such as bone marrow, skin, cartilage, tendons, bone, muscle (including cardiac muscle), blood vessels, cornea, neural cells, gastrointestinal cells and others; and organs for transplantation such as kidney, liver, pancreas (including islet cells), heart and lung.

It is contemplated that multiple different exogenous growth factors and/or cytokines may be administered in combination with the polypeptide of the invention to achieve the desired effect, including any of the growth factors listed herein, other stem cell maintenance factors, and specifically including stem cell factor (SCF), leukemia inhibitory factor (LIF), Flt-3 ligand (Flt-3L), any of the interleukins, recombinant soluble IL-6 receptor fused to IL-6, macrophage inflammatory protein 1-alpha (MIP-1-alpha), G-CSF, GM-CSF, thrombopoietin (TPO), platelet factor 4 (PF-4), platelet-derived growth factor (PDGF), neural growth factors and basic fibroblast growth factor (bFGF).

Since totipotent stem cells can give rise to virtually any mature cell type, expansion of these cells in culture will facilitate the production of large quantities of mature cells. Techniques for culturing stem cells are known in the art and administration of polypeptides of the invention, optionally with other growth factors and/or cytokines, is expected to enhance the survival and proliferation of the stem cell populations. This can be accomplished by direct administration of the polypeptide of the invention to the culture medium. Alternatively, stroma cells transfected with a polynucleotide that encodes for the polypeptide of the invention can be used as a feeder

layer for the stem cell populations in culture or in vivo. Stromal support cells for feeder layers may include embryonic bone marrow fibroblasts, bone marrow stromal cells, fetal liver cells, or cultured embryonic fibroblasts (see U.S. Patent No. 5,690,926).

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Stem cells themselves can be transfected with a polynucleotide of the invention to induce autocrine expression of the polypeptide of the invention. This will allow for generation of undifferentiated totipotential/pluripotential stem cell lines that are useful as is or that can then be differentiated into the desired mature cell types. These stable cell lines can also serve as a source of undifferentiated totipotential/pluripotential mRNA to create cDNA libraries and templates for polymerase chain reaction experiments. These studies would allow for the isolation and identification of differentially expressed genes in stem cell populations that regulate stem cell proliferation and/or maintenance.

Expansion and maintenance of totipotent stem cell populations will be useful in the treatment of many pathological conditions. For example, polypeptides of the present invention may be used to manipulate stem cells in culture to give rise to neuroepithelial cells that can be used to augment or replace cells damaged by illness, autoimmune disease, accidental damage or genetic disorders. The polypeptide of the invention may be useful for inducing the proliferation of neural cells and for the regeneration of nerve and brain tissue, *i.e.* for the treatment of central and peripheral nervous system diseases and neuropathies, as well as mechanical and traumatic disorders which involve degeneration, death or trauma to neural cells or nerve tissue. In addition, the expanded stem cell populations can also be genetically altered for gene therapy purposes and to decrease host rejection of replacement tissues after grafting or implantation.

Expression of the polypeptide of the invention and its effect on stem cells can also be manipulated to achieve controlled differentiation of the stem cells into more differentiated cell types. A broadly applicable method of obtaining pure populations of a specific differentiated cell type from undifferentiated stem cell populations involves the use of a cell-type specific promoter driving a selectable marker. The selectable marker allows only cells of the desired type to survive. For example, stem cells can be induced to differentiate into cardiomyocytes (Wobus et al., Differentiation, 48: 173-182, (1991); Klug et al., J. Clin. Invest., 98(1): 216-224, (1998)) or skeletal muscle cells (Browder, L. W. In: *Principles of Tissue Engineering eds*. Lanza et al., Academic Press (1997)). Alternatively, directed differentiation of stem cells can be accomplished by culturing the stem cells in the presence of a differentiation factor such as retinoic acid and an antagonist of the polypeptide of the invention which would inhibit the effects of endogenous stem cell factor activity and allow differentiation to proceed.

*In vitro* cultures of stem cells can be used to determine if the polypeptide of the invention exhibits stem cell growth factor activity. Stem cells are isolated from any one of various cell

sources (including hematopoietic stem cells and embryonic stem cells) and cultured on a feeder layer, as described by Thompson et al. Proc. Natl. Acad. Sci, U.S.A., 92: 7844-7848 (1995), in the presence of the polypeptide of the invention alone or in combination with other growth factors or cytokines. The ability of the polypeptide of the invention to induce stem cells proliferation is determined by colony formation on semi-solid support *e.g.* as described by Bernstein et al., Blood, 77: 2316-2321 (1991).

#### 4.10.5 HEMATOPOIESIS REGULATING ACTIVITY

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A polypeptide of the present invention may be involved in regulation of hematopoiesis and, consequently, in the treatment of myeloid or lymphoid cell disorders. Even marginal biological activity in support of colony forming cells or of factor-dependent cell lines indicates involvement in regulating hematopoiesis, e.g. in supporting the growth and proliferation of erythroid progenitor cells alone or in combination with other cytokines, thereby indicating utility, for example, in treating various anemias or for use in conjunction with irradiation/chemotherapy to stimulate the production of erythroid precursors and/or erythroid cells; in supporting the growth and proliferation of myeloid cells such as granulocytes and monocytes/macrophages (i.e., traditional CSF activity) useful, for example, in conjunction with chemotherapy to prevent or treat consequent myelo-suppression; in supporting the growth and proliferation of megakaryocytes and consequently of platelets thereby allowing prevention or treatment of various platelet disorders such as thrombocytopenia, and generally for use in place of or complimentary to platelet transfusions; and/or in supporting the growth and proliferation of hematopoietic stem cells which are capable of maturing to any and all of the above-mentioned hematopoietic cells and therefore find therapeutic utility in various stem cell disorders (such as those usually treated with transplantation, including, without limitation, aplastic anemia and paroxysmal nocturnal hemoglobinuria), as well as in repopulating the stem cell compartment post irradiation/chemotherapy, either in-vivo or ex-vivo (i.e., in conjunction with bone marrow transplantation or with peripheral progenitor cell transplantation (homologous or heterologous)) as normal cells or genetically manipulated for gene therapy.

Therapeutic compositions of the invention can be used in the following:

Suitable assays for proliferation and differentiation of various hematopoietic lines are cited above.

Assays for embryonic stem cell differentiation (which will identify, among others, proteins that influence embryonic differentiation hematopoiesis) include, without limitation, those described in: Johansson et al. Cellular Biology 15:141-151, 1995; Keller et al., Molecular and Cellular Biology 13:473-486, 1993; McClanahan et al., Blood 81:2903-2915, 1993.

Assays for stem cell survival and differentiation (which will identify, among others, proteins that regulate lympho-hematopoiesis) include, without limitation, those described in: Methylcellulose colony forming assays, Freshney, M. G. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 265-268, Wiley-Liss, Inc., New York, N.Y. 1994; Hirayama et al., Proc. Natl. Acad. Sci. USA 89:5907-5911, 1992; Primitive hematopoietic colony forming cells with high proliferative potential, McNiece, I. K. and Briddell, R. A. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 23-39, Wiley-Liss, Inc., New York, N.Y. 1994; Neben et al., Experimental Hematology 22:353-359, 1994; Cobblestone area forming cell assay, Ploemacher, R. E. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 1-21, Wiley-Liss, Inc., New York, N.Y. 1994; Long term bone marrow cultures in the presence of stromal cells, Spooncer, E., Dexter, M. and Allen, T. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 163-179, Wiley-Liss, Inc., New York, N.Y. 1994; Long term culture initiating cell assay, Sutherland, H. J. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 139-162, Wiley-Liss, Inc., New York, N.Y. 1994.

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## 4.10.6 TISSUE GROWTH ACTIVITY

A polypeptide of the present invention also may be involved in bone, cartilage, tendon, ligament and/or nerve tissue growth or regeneration, as well as in wound healing and tissue repair and replacement, and in healing of burns, incisions and ulcers.

A polypeptide of the present invention which induces cartilage and/or bone growth in circumstances where bone is not normally formed, has application in the healing of bone fractures and cartilage damage or defects in humans and other animals. Compositions of a polypeptide, antibody, binding partner, or other modulator of the invention may have prophylactic use in closed as well as open fracture reduction and also in the improved fixation of artificial joints. De novo bone formation induced by an osteogenic agent contributes to the repair of congenital, trauma induced, or oncologic resection induced craniofacial defects, and also is useful in cosmetic plastic surgery.

A polypeptide of this invention may also be involved in attracting bone-forming cells, stimulating growth of bone-forming cells, or inducing differentiation of progenitors of bone-forming cells. Treatment of osteoporosis, osteoarthritis, bone degenerative disorders, or periodontal disease, such as through stimulation of bone and/or cartilage repair or by blocking inflammation or processes of tissue destruction (collagenase activity, osteoclast activity, etc.) mediated by inflammatory processes may also be possible using the composition of the invention.

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Another category of tissue regeneration activity that may involve the polypeptide of the present invention is tendon/ligament formation. Induction of tendon/ligament-like tissue or other tissue formation in circumstances where such tissue is not normally formed, has application in the healing of tendon or ligament tears, deformities and other tendon or ligament defects in humans and other animals. Such a preparation employing a tendon/ligament-like tissue inducing protein may have prophylactic use in preventing damage to tendon or ligament tissue, as well as use in the improved fixation of tendon or ligament to bone or other tissues, and in repairing defects to tendon or ligament tissue. De novo tendon/ligament-like tissue formation induced by a composition of the present invention contributes to the repair of congenital, trauma induced, or other tendon or ligament defects of other origin, and is also useful in cosmetic plastic surgery for attachment or repair of tendons or ligaments. The compositions of the present invention may provide environment to attract tendon- or ligament-forming cells, stimulate growth of tendon- or ligament-forming cells, induce differentiation of progenitors of tendon- or ligament-forming cells, or induce growth of tendon/ligament cells or progenitors ex vivo for return in vivo to effect tissue repair. The compositions of the invention may also be useful in the treatment of tendinitis, carpal tunnel syndrome and other tendon or ligament defects. The compositions may also include an appropriate matrix and/or sequestering agent as a carrier as is well known in the art.

The compositions of the present invention may also be useful for proliferation of neural cells and for regeneration of nerve and brain tissue, *i.e.* for the treatment of central and peripheral nervous system diseases and neuropathies, as well as mechanical and traumatic disorders, which involve degeneration, death or trauma to neural cells or nerve tissue. More specifically, a composition may be used in the treatment of diseases of the peripheral nervous system, such as peripheral nerve injuries, peripheral neuropathy and localized neuropathies, and central nervous system diseases, such as Alzheimer's, Parkinson's disease, Huntington's disease, amyotrophic lateral sclerosis, and Shy-Drager syndrome. Further conditions which may be treated in accordance with the present invention include mechanical and traumatic disorders, such as spinal cord disorders, head trauma and cerebrovascular diseases such as stroke. Peripheral neuropathies resulting from chemotherapy or other medical therapies may also be treatable using a composition of the invention.

Compositions of the invention may also be useful to promote better or faster closure of non-healing wounds, including without limitation pressure ulcers, ulcers associated with vascular insufficiency, surgical and traumatic wounds, and the like.

Compositions of the present invention may also be involved in the generation or regeneration of other tissues, such as organs (including, for example, pancreas, liver, intestine,

kidney, skin, endothelium), muscle (smooth, skeletal or cardiac) and vascular (including vascular endothelium) tissue, or for promoting the growth of cells comprising such tissues. Part of the desired effects may be by inhibition or modulation of fibrotic scarring may allow normal tissue to regenerate. A polypeptide of the present invention may also exhibit angiogenic activity.

A composition of the present invention may also be useful for gut protection or regeneration and treatment of lung or liver fibrosis, reperfusion injury in various tissues, and conditions resulting from systemic cytokine damage.

A composition of the present invention may also be useful for promoting or inhibiting differentiation of tissues described above from precursor tissues or cells; or for inhibiting the growth of tissues described above.

Therapeutic compositions of the invention can be used in the following:

Assays for tissue generation activity include, without limitation, those described in: International Patent Publication No. WO95/16035 (bone, cartilage, tendon); International Patent Publication No. WO95/05846 (nerve, neuronal); International Patent Publication No. WO91/07491 (skin, endothelium).

Assays for wound healing activity include, without limitation, those described in: Winter, Epidermal Wound Healing, pps. 71-112 (Maibach, H. I. and Rovee, D. T., eds.), Year Book Medical Publishers, Inc., Chicago, as modified by Eaglstein and Mertz, J. Invest. Dermatol 71:382-84 (1978).

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## 4.10.7 IMMUNE STIMULATING OR SUPPRESSING ACTIVITY

A polypeptide of the present invention may also exhibit immune stimulating or immune suppressing activity, including without limitation the activities for which assays are described herein. A polynucleotide of the invention can encode a polypeptide exhibiting such activities. A protein may be useful in the treatment of various immune deficiencies and disorders (including severe combined immunodeficiency (SCID)), *e.g.*, in regulating (up or down) growth and proliferation of T and/or B lymphocytes, as well as effecting the cytolytic activity of NK cells and other cell populations. These immune deficiencies may be genetic or be caused by viral (*e.g.*, HIV) as well as bacterial or fungal infections, or may result from autoimmune disorders. More specifically, infectious diseases causes by viral, bacterial, fungal or other infection may be treatable using a protein of the present invention, including infections by HIV, hepatitis viruses, herpes viruses, mycobacteria, Leishmania spp., malaria spp. and various fungal infections such as candidiasis. Of course, in this regard, proteins of the present invention may also be useful where a boost to the immune system generally may be desirable, *i.e.*, in the treatment of cancer.

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Autoimmune disorders which may be treated using a protein of the present invention include, for example, connective tissue disease, multiple sclerosis, systemic lupus erythematosus, rheumatoid arthritis, autoimmune pulmonary inflammation, Guillain-Barre syndrome, autoimmune thyroiditis, insulin dependent diabetes mellitis, myasthenia gravis, graft-versus-host disease and autoimmune inflammatory eye disease. Such a protein (or antagonists thereof, including antibodies) of the present invention may also to be useful in the treatment of allergic reactions and conditions (e.g., anaphylaxis, serum sickness, drug reactions, food allergies, insect venom allergies, mastocytosis, allergic rhinitis, hypersensitivity pneumonitis, urticaria, angioedema, eczema, atopic dermatitis, allergic contact dermatitis, erythema multiforme, Stevens-Johnson syndrome, allergic conjunctivitis, atopic keratoconjunctivitis, venereal keratoconjunctivitis, giant papillary conjunctivitis and contact allergies), such as asthma (particularly allergic asthma) or other respiratory problems. Other conditions, in which immune suppression is desired (including, for example, organ transplantation), may also be treatable using a protein (or antagonists thereof) of the present invention. The therapeutic effects of the polypeptides or antagonists thereof on allergic reactions can be evaluated by in vivo animals models such as the cumulative contact enhancement test (Lastborn et al., Toxicology 125: 59-66, 1998), skin prick test (Hoffmann et al., Allergy 54: 446-54, 1999), guinea pig skin sensitization test (Vohr et al., Arch. Toxocol. 73: 501-9), and murine local lymph node assay (Kimber et al., J. Toxicol. Environ. Health 53: 563-79).

Using the proteins of the invention it may also be possible to modulate immune responses, in a number of ways. Down regulation may be in the form of inhibiting or blocking an immune response already in progress or may involve preventing the induction of an immune response. The functions of activated T cells may be inhibited by suppressing T cell responses or by inducing specific tolerance in T cells, or both. Immunosuppression of T cell responses is generally an active, non-antigen-specific, process which requires continuous exposure of the T cells to the suppressive agent. Tolerance, which involves inducing non-responsiveness or anergy in T cells, is distinguishable from immunosuppression in that it is generally antigen-specific and persists after exposure to the tolerizing agent has ceased. Operationally, tolerance can be demonstrated by the lack of a T cell response upon reexposure to specific antigen in the absence of the tolerizing agent.

Down regulating or preventing one or more antigen functions (including without limitation B lymphocyte antigen functions (such as, for example, B7)), *e.g.*, preventing high level lymphokine synthesis by activated T cells, will be useful in situations of tissue, skin and organ transplantation and in graft-versus-host disease (GVHD). For example, blockage of T cell function should result in reduced tissue destruction in tissue transplantation. Typically, in tissue

transplants, rejection of the transplant is initiated through its recognition as foreign by T cells, followed by an immune reaction that destroys the transplant. The administration of a therapeutic composition of the invention may prevent cytokine synthesis by immune cells, such as T cells, and thus acts as an immunosuppressant. Moreover, a lack of costimulation may also be sufficient to anergize the T cells, thereby inducing tolerance in a subject. Induction of long-term tolerance by B lymphocyte antigen-blocking reagents may avoid the necessity of repeated administration of these blocking reagents. To achieve sufficient immunosuppression or tolerance in a subject, it may also be necessary to block the function of a combination of B lymphocyte antigens.

The efficacy of particular therapeutic compositions in preventing organ transplant rejection or GVHD can be assessed using animal models that are predictive of efficacy in humans. Examples of appropriate systems which can be used include allogeneic cardiac grafts in rats and xenogeneic pancreatic islet cell grafts in mice, both of which have been used to examine the immunosuppressive effects of CTLA4Ig fusion proteins in vivo as described in Lenschow et al., Science 257:789-792 (1992) and Turka et al., Proc. Natl. Acad. Sci USA, 89:11102-11105 (1992). In addition, murine models of GVHD (see Paul ed., Fundamental Immunology, Raven Press, New York, 1989, pp. 846-847) can be used to determine the effect of therapeutic compositions of the invention on the development of that disease.

Blocking antigen function may also be therapeutically useful for treating autoimmune diseases. Many autoimmune disorders are the result of inappropriate activation of T cells that are reactive against self tissue and which promote the production of cytokines and autoantibodies involved in the pathology of the diseases. Preventing the activation of autoreactive T cells may reduce or eliminate disease symptoms. Administration of reagents which block stimulation of T cells can be used to inhibit T cell activation and prevent production of autoantibodies or T cell-derived cytokines which may be involved in the disease process. Additionally, blocking reagents may induce antigen-specific tolerance of autoreactive T cells which could lead to long-term relief from the disease. The efficacy of blocking reagents in preventing or alleviating autoimmune disorders can be determined using a number of well-characterized animal models of human autoimmune diseases. Examples include murine experimental autoimmune encephalitis, systemic lupus erythmatosis in MRL/lpr/lpr mice or NZB hybrid mice, murine autoimmune collagen arthritis, diabetes mellitus in NOD mice and BB rats, and murine experimental myasthenia gravis (see Paul ed., Fundamental Immunology, Raven Press, New York, 1989, pp. 840-856).

Upregulation of an antigen function (e.g., a B lymphocyte antigen function), as a means of up regulating immune responses, may also be useful in therapy. Upregulation of immune responses may be in the form of enhancing an existing immune response or eliciting an initial

immune response. For example, enhancing an immune response may be useful in cases of viral infection, including systemic viral diseases such as influenza, the common cold, and encephalitis.

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Alternatively, anti-viral immune responses may be enhanced in an infected patient by removing T cells from the patient, costimulating the T cells in vitro with viral antigen-pulsed APCs either expressing a peptide of the present invention or together with a stimulatory form of a soluble peptide of the present invention and reintroducing the in vitro activated T cells into the patient. Another method of enhancing anti-viral immune responses would be to isolate infected cells from a patient, transfect them with a nucleic acid encoding a protein of the present invention as described herein such that the cells express all or a portion of the protein on their surface, and reintroduce the transfected cells into the patient. The infected cells would now be capable of delivering a costimulatory signal to, and thereby activate, T cells in vivo.

A polypeptide of the present invention may provide the necessary stimulation signal to T cells to induce a T cell mediated immune response against the transfected tumor cells. In addition, tumor cells which lack MHC class I or MHC class II molecules, or which fail to reexpress sufficient mounts of MHC class I or MHC class II molecules, can be transfected with nucleic acid encoding all or a portion of (e.g., a cytoplasmic-domain truncated portion) of an MHC class I alpha chain protein and β<sub>2</sub> microglobulin protein or an MHC class II alpha chain protein and an MHC class II beta chain protein to thereby express MHC class I or MHC class II proteins on the cell surface. Expression of the appropriate class I or class II MHC in conjunction with a peptide having the activity of a B lymphocyte antigen (e.g., B7-1, B7-2, B7-3) induces a T cell mediated immune response against the transfected tumor cell. Optionally, a gene encoding an antisense construct which blocks expression of an MHC class II associated protein, such as the invariant chain, can also be cotransfected with a DNA encoding a peptide having the activity of a B lymphocyte antigen to promote presentation of tumor associated antigens and induce tumor specific immunity. Thus, the induction of a T cell mediated immune response in a human subject may be sufficient to overcome tumor-specific tolerance in the subject.

The activity of a protein of the invention may, among other means, be measured by the following methods:

Suitable assays for thymocyte or splenocyte cytotoxicity include, without limitation, those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, In Vitro assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Herrmann et al., Proc. Natl. Acad. Sci. USA 78:2488-2492, 1981; Herrmann et al., J. Immunol. 128:1968-1974, 1982; Handa et al., J.

Immunol. 135:1564-1572, 1985; Takai et al., I. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988; Bowman et al., J. Virology 61:1992-1998; Bertagnolli et al., Cellular Immunology 133:327-341, 1991; Brown et al., J. Immunol. 153:3079-3092, 1994.

Assays for T-cell-dependent immunoglobulin responses and isotype switching (which will identify, among others, proteins that modulate T-cell dependent antibody responses and that affect Th1/Th2 profiles) include, without limitation, those described in: Maliszewski, J. Immunol. 144:3028-3033, 1990; and Assays for B cell function: In vitro antibody production, Mond, J. J. and Brunswick, M. In Current Protocols in Immunology. J. E. e.a. Coligan eds. Vol 1 pp. 3.8.1-3.8.16, John Wiley and Sons, Toronto. 1994.

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Mixed lymphocyte reaction (MLR) assays (which will identify, among others, proteins that generate predominantly Th1 and CTL responses) include, without limitation, those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, In Vitro assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988; Bertagnolli et al., J. Immunol. 149:3778-3783, 1992.

Dendritic cell-dependent assays (which will identify, among others, proteins expressed by dendritic cells that activate naive T-cells) include, without limitation, those described in: Guery et al., J. Immunol. 134:536-544, 1995; Inaba et al., Journal of Experimental Medicine 173:549-559, 1991; Macatonia et al., Journal of Immunology 154:5071-5079, 1995; Porgador et al., Journal of Experimental Medicine 182:255-260, 1995; Nair et al., Journal of Virology 67:4062-4069, 1993; Huang et al., Science 264:961-965, 1994; Macatonia et al., Journal of Experimental Medicine 169:1255-1264, 1989; Bhardwaj et al., Journal of Clinical Investigation 94:797-807, 1994; and Inaba et al., Journal of Experimental Medicine 172:631-640, 1990.

Assays for lymphocyte survival/apoptosis (which will identify, among others, proteins that prevent apoptosis after superantigen induction and proteins that regulate lymphocyte homeostasis) include, without limitation, those described in: Darzynkiewicz et al., Cytometry 13:795-808, 1992; Gorczyca et al., Leukemia 7:659-670, 1993; Gorczyca et al., Cancer Research 53:1945-1951, 1993; Itoh et al., Cell 66:233-243, 1991; Zacharchuk, Journal of Immunology 145:4037-4045, 1990; Zamai et al., Cytometry 14:891-897, 1993; Gorczyca et al., International Journal of Oncology 1:639-648, 1992.

Assays for proteins that influence early steps of T-cell commitment and development include, without limitation, those described in: Antica et al., Blood 84:111-117, 1994; Fine et al., Cellular Immunology 155:111-122, 1994; Galy et al., Blood 85:2770-2778, 1995; Toki et al., Proc. Nat. Acad Sci. USA 88:7548-7551, 1991.

## 4.10.8 ACTIVIN/INHIBIN ACTIVITY

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A polypeptide of the present invention may also exhibit activin- or inhibin-related activities. A polynucleotide of the invention may encode a polypeptide exhibiting such characteristics. Inhibins are characterized by their ability to inhibit the release of follicle stimulating hormone (FSH), while activins and are characterized by their ability to stimulate the release of follicle stimulating hormone (FSH). Thus, a polypeptide of the present invention, alone or in heterodimers with a member of the inhibin family, may be useful as a contraceptive based on the ability of inhibins to decrease fertility in female mammals and decrease spermatogenesis in male mammals. Administration of sufficient amounts of other inhibins can induce infertility in these mammals. Alternatively, the polypeptide of the invention, as a homodimer or as a heterodimer with other protein subunits of the inhibin group, may be useful as a fertility inducing therapeutic, based upon the ability of activin molecules in stimulating FSH release from cells of the anterior pituitary. See, for example, U.S. Pat. No. 4,798,885. A polypeptide of the invention may also be useful for advancement of the onset of fertility in sexually immature mammals, so as to increase the lifetime reproductive performance of domestic animals such as, but not limited to, cows, sheep and pigs.

The activity of a polypeptide of the invention may, among other means, be measured by the following methods.

Assays for activin/inhibin activity include, without limitation, those described in: Vale et al., Endocrinology 91:562-572, 1972; Ling et al., Nature 321:779-782, 1986; Vale et al., Nature 321:776-779, 1986; Mason et al., Nature 318:659-663, 1985; Forage et al., Proc. Natl. Acad. Sci. USA 83:3091-3095, 1986.

## 4.10.9 CHEMOTACTIC/CHEMOKINETIC ACTIVITY

A polypeptide of the present invention may be involved in chemotactic or chemokinetic activity for mammalian cells, including, for example, monocytes, fibroblasts, neutrophils, T-cells, mast cells, eosinophils, epithelial and/or endothelial cells. A polynucleotide of the invention can encode a polypeptide exhibiting such attributes. Chemotactic and chemokinetic receptor activation can be used to mobilize or attract a desired cell population to a desired site of action. Chemotactic or chemokinetic compositions (e.g. proteins, antibodies, binding partners, or modulators of the invention) provide particular advantages in treatment of wounds and other trauma to tissues, as well as in treatment of localized infections. For example, attraction of lymphocytes, monocytes or neutrophils to tumors or sites of infection may result in improved immune responses against the tumor or infecting agent.

A protein or peptide has chemotactic activity for a particular cell population if it can stimulate, directly or indirectly, the directed orientation or movement of such cell population. Preferably, the protein or peptide has the ability to directly stimulate directed movement of cells. Whether a particular protein has chemotactic activity for a population of cells can be readily determined by employing such protein or peptide in any known assay for cell chemotaxis.

Therapeutic compositions of the invention can be used in the following:

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Assays for chemotactic activity (which will identify proteins that induce or prevent chemotaxis) consist of assays that measure the ability of a protein to induce the migration of cells across a membrane as well as the ability of a protein to induce the adhesion of one cell population to another cell population. Suitable assays for movement and adhesion include, without limitation, those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Marguiles, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 6.12, Measurement of alpha and beta Chemokines 6.12.1-6.12.28; Taub et al. J. Clin. Invest. 95:1370-1376, 1995; Lind et al. APMIS 103:140-146, 1995; Muller et al Eur. J. Immunol. 25:1744-1748; Gruber et al. J. of Immunol. 152:5860-5867, 1994; Johnston et al. J. of Immunol. 153:1762-1768, 1994.

## 4.10.10 HEMOSTATIC AND THROMBOLYTIC ACTIVITY

A polypeptide of the invention may also be involved in hemostatis or thrombolysis or thrombosis. A polynucleotide of the invention can encode a polypeptide exhibiting such attributes. Compositions may be useful in treatment of various coagulation disorders (including hereditary disorders, such as hemophilias) or to enhance coagulation and other hemostatic events in treating wounds resulting from trauma, surgery or other causes. A composition of the invention may also be useful for dissolving or inhibiting formation of thromboses and for treatment and prevention of conditions resulting therefrom (such as, for example, infarction of cardiac and central nervous system vessels (e.g., stroke).

Therapeutic compositions of the invention can be used in the following:

Assay for hemostatic and thrombolytic activity include, without limitation, those described in: Linet et al., J. Clin. Pharmacol. 26:131-140, 1986; Burdick et al., Thrombosis Res. 45:413-419, 1987; Humphrey et al., Fibrinolysis 5:71-79 (1991); Schaub, Prostaglandins 35:467-474, 1988.

## 4.10.11 CANCER DIAGNOSIS AND THERAPY

Polypeptides of the invention may be involved in cancer cell generation, proliferation or metastasis. Detection of the presence or amount of polynucleotides or polypeptides of the

invention may be useful for the diagnosis and/or prognosis of one or more types of cancer. For example, the presence or increased expression of a polynucleotide/polypeptide of the invention may indicate a hereditary risk of cancer, a precancerous condition, or an ongoing malignancy. Conversely, a defect in the gene or absence of the polypeptide may be associated with a cancer condition. Identification of single nucleotide polymorphisms associated with cancer or a predisposition to cancer may also be useful for diagnosis or prognosis.

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Cancer treatments promote tumor regression by inhibiting tumor cell proliferation, inhibiting angiogenesis (growth of new blood vessels that is necessary to support tumor growth) and/or prohibiting metastasis by reducing tumor cell motility or invasiveness. Therapeutic compositions of the invention may be effective in adult and pediatric oncology including in solid phase tumors/malignancies, locally advanced tumors, human soft tissue sarcomas, metastatic cancer, including lymphatic metastases, blood cell malignancies including multiple myeloma, acute and chronic leukemias, and lymphomas, head and neck cancers including mouth cancer, larynx cancer and thyroid cancer, lung cancers including small cell carcinoma and non-small cell cancers, breast cancers including small cell carcinoma and ductal carcinoma, gastrointestinal cancers including esophageal cancer, stomach cancer, colon cancer, colorectal cancer and polyps associated with colorectal neoplasia, pancreatic cancers, liver cancer, urologic cancers including bladder cancer and prostate cancer, malignancies of the female genital tract including ovarian carcinoma, uterine (including endometrial) cancers, and solid tumor in the ovarian follicle, kidney cancers including renal cell carcinoma, brain cancers including intrinsic brain tumors, neuroblastoma, astrocytic brain tumors, gliomas, metastatic tumor cell invasion in the central nervous system, bone cancers including osteomas, skin cancers including malignant melanoma, tumor progression of human skin keratinocytes, squamous cell carcinoma, basal cell carcinoma, hemangiopericytoma and Karposi's sarcoma.

Polypeptides, polynucleotides, or modulators of polypeptides of the invention (including inhibitors and stimulators of the biological activity of the polypeptide of the invention) may be administered to treat cancer. Therapeutic compositions can be administered in therapeutically effective dosages alone or in combination with adjuvant cancer therapy such as surgery, chemotherapy, radiotherapy, thermotherapy, and laser therapy, and may provide a beneficial effect, *e.g.* reducing tumor size, slowing rate of tumor growth, inhibiting metastasis, or otherwise improving overall clinical condition, without necessarily eradicating the cancer.

The composition can also be administered in therapeutically effective amounts as a portion of an anti-cancer cocktail. An anti-cancer cocktail is a mixture of the polypeptide or modulator of the invention with one or more anti-cancer drugs in addition to a pharmaceutically acceptable carrier for delivery. The use of anti-cancer cocktails as a cancer treatment is routine.

Anti-cancer drugs that are well known in the art and can be used as a treatment in combination with the polypeptide or modulator of the invention include: Actinomycin D, Aminoglutethimide, Asparaginase, Bleomycin, Busulfan, Carboplatin, Carmustine, Chlorambucil, Cisplatin (cis-DDP), Cyclophosphamide, Cytarabine HCl (Cytosine arabinoside), Dacarbazine, Dactinomycin, Daunorubicin HCl, Doxorubicin HCl, Estramustine phosphate sodium, Etoposide (V16-213), Floxuridine, 5-Fluorouracil (5-Fu), Flutamide, Hydroxyurea (hydroxycarbamide), Ifosfamide, Interferon Alpha-2a, Interferon Alpha-2b, Leuprolide acetate (LHRH-releasing factor analog), Lomustine, Mechlorethamine HCl (nitrogen°mustard), Melphalan, Mercaptopurine, Mesna, Methotrexate (MTX), Mitomycin, Mitoxantrone HCl, Octreotide, Plicamycin, Procarbazine HCl, Streptozocin, Tamoxifen citrate, Thioguanine, Thiotepa, Vinblastine sulfate, Vincristine sulfate, Amsacrine, Azacitidine, Hexamethylmelamine, Interleukin-2, Mitoguazone, Pentostatin, Semustine, Teniposide, and Vindesine sulfate.

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In addition, therapeutic compositions of the invention may be used for prophylactic treatment of cancer. There are hereditary conditions and/or environmental situations (e.g. exposure to carcinogens) known in the art that predispose an individual to developing cancers. Under these circumstances, it may be beneficial to treat these individuals with therapeutically effective doses of the polypeptide of the invention to reduce the risk of developing cancers.

In vitro models can be used to determine the effective doses of the polypeptide of the invention as a potential cancer treatment. These *in vitro* models include proliferation assays of cultured tumor cells, growth of cultured tumor cells in soft agar (see Freshney, (1987) Culture of Animal Cells: A Manual of Basic Technique, Wily-Liss, New York, NY Ch 18 and Ch 21), tumor systems in nude mice as described in Giovanella et al., J. Natl. Can. Inst., 52: 921-30 (1974), mobility and invasive potential of tumor cells in Boyden Chamber assays as described in Pilkington et al., Anticancer Res., 17: 4107-9 (1997), and angiogenesis assays such as induction of vascularization of the chick chorioallantoic membrane or induction of vascular endothelial cell migration as described in Ribatta et al., Intl. J. Dev. Biol., 40: 1189-97 (1999) and Li et al., Clin. Exp. Metastasis, 17:423-9 (1999), respectively. Suitable tumor cells lines are available, *e.g.* from American Type Tissue Culture Collection catalogs.

# 4.10.12 RECEPTOR/LIGAND ACTIVITY

A polypeptide of the present invention may also demonstrate activity as receptor, receptor ligand or inhibitor or agonist of receptor/ligand interactions. A polynucleotide of the invention can encode a polypeptide exhibiting such characteristics. Examples of such receptors and ligands include, without limitation, cytokine receptors and their ligands, receptor kinases and their ligands, receptor phosphatases and their ligands, receptors involved in cell-cell interactions

and their ligands (including without limitation, cellular adhesion molecules (such as selectins, integrins and their ligands) and receptor/ligand pairs involved in antigen presentation, antigen recognition and development of cellular and humoral immune responses. Receptors and ligands are also useful for screening of potential peptide or small molecule inhibitors of the relevant receptor/ligand interaction. A protein of the present invention (including, without limitation, fragments of receptors and ligands) may themselves be useful as inhibitors of receptor/ligand interactions.

The activity of a polypeptide of the invention may, among other means, be measured by the following methods:

Suitable assays for receptor-ligand activity include without limitation those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley- Interscience (Chapter 7.28, Measurement of Cellular Adhesion under static conditions 7.28.1-7.28.22), Takai et al., Proc. Natl. Acad. Sci. USA 84:6864-6868, 1987; Bierer et al., J. Exp. Med. 168:1145-1156, 1988; Rosenstein et al., J. Exp. Med. 169:149-160 1989; Stoltenborg et al., J. Immunol. Methods 175:59-68, 1994; Stitt et al., Cell 80:661-670, 1995.

By way of example, the polypeptides of the invention may be used as a receptor for a ligand(s) thereby transmitting the biological activity of that ligand(s). Ligands may be identified through binding assays, affinity chromatography, dihybrid screening assays, BIAcore assays, gel overlay assays, or other methods known in the art.

Studies characterizing drugs or proteins as agonist or antagonist or partial agonists or a partial antagonist require the use of other proteins as competing ligands. The polypeptides of the present invention or ligand(s) thereof may be labeled by being coupled to radioisotopes, colorimetric molecules or a toxin molecules by conventional methods. ("Guide to Protein Purification" Murray P. Deutscher (ed) Methods in Enzymology Vol. 182 (1990) Academic Press, Inc. San Diego). Examples of radioisotopes include, but are not limited to, tritium and carbon-14. Examples of colorimetric molecules include, but are not limited to, fluorescent molecules such as fluorescamine, or rhodamine or other colorimetric molecules. Examples of toxins include, but are not limited, to ricin.

#### 4.10.13 DRUG SCREENING

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This invention is particularly useful for screening chemical compounds by using the novel polypeptides or binding fragments thereof in any of a variety of drug screening techniques. The polypeptides or fragments employed in such a test may either be free in solution, affixed to a solid support, borne on a cell surface or located intracellularly. One method of drug screening

utilizes eukaryotic or prokaryotic host cells which are stably transformed with recombinant nucleic acids expressing the polypeptide or a fragment thereof. Drugs are screened against such transformed cells in competitive binding assays. Such cells, either in viable or fixed form, can be used for standard binding assays. One may measure, for example, the formation of complexes between polypeptides of the invention or fragments and the agent being tested or examine the diminution in complex formation between the novel polypeptides and an appropriate cell line, which are well known in the art.

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Sources for test compounds that may be screened for ability to bind to or modulate (*i.e.*, increase or decrease) the activity of polypeptides of the invention include (1) inorganic and organic chemical libraries, (2) natural product libraries, and (3) combinatorial libraries comprised of either random or mimetic peptides, oligonucleotides or organic molecules.

Chemical libraries may be readily synthesized or purchased from a number of commercial sources, and may include structural analogs of known compounds or compounds that are identified as "hits" or "leads" via natural product screening.

The sources of natural product libraries are microorganisms (including bacteria and fungi), animals, plants or other vegetation, or marine organisms, and libraries of mixtures for screening may be created by: (1) fermentation and extraction of broths from soil, plant or marine microorganisms or (2) extraction of the organisms themselves. Natural product libraries include polyketides, non-ribosomal peptides, and (non-naturally occurring) variants thereof. For a review, see *Science 282*:63-68 (1998).

Combinatorial libraries are composed of large numbers of peptides, oligonucleotides or organic compounds and can be readily prepared by traditional automated synthesis methods, PCR, cloning or proprietary synthetic methods. Of particular interest are peptide and oligonucleotide combinatorial libraries. Still other libraries of interest include peptide, protein, peptidomimetic, multiparallel synthetic collection, recombinatorial, and polypeptide libraries. For a review of combinatorial chemistry and libraries created therefrom, see Myers, *Curr. Opin. Biotechnol.* 8:701-707 (1997). For reviews and examples of peptidomimetic libraries, see Al-Obeidi et al., *Mol. Biotechnol.* 9(3):205-23 (1998); Hruby et al., *Curr Opin Chem Biol.* 1(1):114-19 (1997); Dorner et al., *Bioorg Med Chem.* 4(5):709-15 (1996) (alkylated dipeptides).

Identification of modulators through use of the various libraries described herein permits modification of the candidate "hit" (or "lead") to optimize the capacity of the "hit" to bind a polypeptide of the invention. The molecules identified in the binding assay are then tested for antagonist or agonist activity in *in vivo* tissue culture or animal models that are well known in the art. In brief, the molecules are titrated into a plurality of cell cultures or animals and then tested for either cell/animal death or prolonged survival of the animal/cells.

The binding molecules thus identified may be complexed with toxins, e.g., ricin or cholera, or with other compounds that are toxic to cells such as radioisotopes. The toxin-binding molecule complex is then targeted to a tumor or other cell by the specificity of the binding molecule for a polypeptide of the invention. Alternatively, the binding molecules may be complexed with imaging agents for targeting and imaging purposes.

# 4.10.14 ASSAY FOR RECEPTOR ACTIVITY

The invention also provides methods to detect specific binding of a polypeptide e.g. a ligand or a receptor. The art provides numerous assays particularly useful for identifying previously unknown binding partners for receptor polypeptides of the invention. For example, expression cloning using mammalian or bacterial cells, or dihybrid screening assays can be used to identify polynucleotides encoding binding partners. As another example, affinity chromatography with the appropriate immobilized polypeptide of the invention can be used to isolate polypeptides that recognize and bind polypeptides of the invention. There are a number of different libraries used for the identification of compounds, and in particular small molecules, that modulate (i.e., increase or decrease) biological activity of a polypeptide of the invention. Ligands for receptor polypeptides of the invention can also be identified by adding exogenous ligands, or cocktails of ligands to two cells populations that are genetically identical except for the expression of the receptor of the invention: one cell population expresses the receptor of the invention whereas the other does not. The response of the two cell populations to the addition of ligands(s) are then compared. Alternatively, an expression library can be co-expressed with the polypeptide of the invention in cells and assayed for an autocrine response to identify potential ligand(s). As still another example, BIAcore assays, gel overlay assays, or other methods known in the art can be used to identify binding partner polypeptides, including, (1) organic and inorganic chemical libraries, (2) natural product libraries, and (3) combinatorial libraries comprised of random peptides, oligonucleotides or organic molecules.

The role of downstream intracellular signaling molecules in the signaling cascade of the polypeptide of the invention can be determined. For example, a chimeric protein in which the cytoplasmic domain of the polypeptide of the invention is fused to the extracellular portion of a protein, whose ligand has been identified, is produced in a host cell. The cell is then incubated with the ligand specific for the extracellular portion of the chimeric protein, thereby activating the chimeric receptor. Known downstream proteins involved in intracellular signaling can then be assayed for expected modifications *i.e.* phosphorylation. Other methods known to those in the art can also be used to identify signaling molecules involved in receptor activity.

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## 4.10.15 ANTI-INFLAMMATORY ACTIVITY

Compositions of the present invention may also exhibit anti-inflammatory activity. The anti-inflammatory activity may be achieved by providing a stimulus to cells involved in the inflammatory response, by inhibiting or promoting cell-cell interactions (such as, for example, cell adhesion), by inhibiting or promoting chemotaxis of cells involved in the inflammatory process, inhibiting or promoting cell extravasation, or by stimulating or suppressing production of other factors which more directly inhibit or promote an inflammatory response. Compositions with such activities can be used to treat inflammatory conditions including chronic or acute conditions), including without limitation intimation associated with infection (such as septic shock, sepsis or systemic inflammatory response syndrome (SIRS)), ischemia-reperfusion injury, endotoxin lethality, arthritis, complement-mediated hyperacute rejection, nephritis, cytokine or chemokine-induced lung injury, inflammatory bowel disease, Crohn's disease or resulting from over production of cytokines such as TNF or IL-1. Compositions of the invention may also be useful to treat anaphylaxis and hypersensitivity to an antigenic substance or material. Compositions of this invention may be utilized to prevent or treat conditions such as, but not limited to, sepsis, acute pancreatitis, endotoxin shock, cytokine induced shock, rheumatoid arthritis, chronic inflammatory arthritis, pancreatic cell damage from diabetes mellitus type 1, graft versus host disease, inflammatory bowel disease, inflamation associated with pulmonary disease, other autoimmune disease or inflammatory disease, an antiproliferative agent such as for acute or chronic mylegenous leukemia or in the prevention of premature labor secondary to intrauterine infections.

#### **4.10.16 LEUKEMIAS**

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Leukemias and related disorders may be treated or prevented by administration of a therapeutic that promotes or inhibits function of the polynucleotides and/or polypeptides of the invention. Such leukemias and related disorders include but are not limited to acute leukemia, acute lymphocytic leukemia, acute myelocytic leukemia, myeloblastic, promyelocytic, myelomonocytic, monocytic, erythroleukemia, chronic leukemia, chronic myelocytic (granulocytic) leukemia and chronic lymphocytic leukemia (for a review of such disorders, see Fishman et al., 1985, Medicine, 2d Ed., J.B. Lippincott Co., Philadelphia).

## 4.10.17 NERVOUS SYSTEM DISORDERS

Nervous system disorders, involving cell types which can be tested for efficacy of intervention with compounds that modulate the activity of the polynucleotides and/or polypeptides of the invention, and which can be treated upon thus observing an indication of

therapeutic utility, include but are not limited to nervous system injuries, and diseases or disorders which result in either a disconnection of axons, a diminution or degeneration of neurons, or demyelination. Nervous system lesions which may be treated in a patient (including human and non-human mammalian patients) according to the invention include but are not limited to the following lesions of either the central (including spinal cord, brain) or peripheral nervous systems:

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- (i) traumatic lesions, including lesions caused by physical injury or associated with surgery, for example, lesions which sever a portion of the nervous system, or compression injuries;
- (ii) ischemic lesions, in which a lack of oxygen in a portion of the nervous system results in neuronal injury or death, including cerebral infarction or ischemia, or spinal cord infarction or ischemia;
- (iii) infectious lesions, in which a portion of the nervous system is destroyed or injured as a result of infection, for example, by an abscess or associated with infection by human immunodeficiency virus, herpes zoster, or herpes simplex virus or with Lyme disease, tuberculosis, syphilis;
- (iv) degenerative lesions, in which a portion of the nervous system is destroyed or injured as a result of a degenerative process including but not limited to degeneration associated with Parkinson's disease, Alzheimer's disease, Huntington's chorea, or amyotrophic lateral sclerosis;
- (v) lesions associated with nutritional diseases or disorders, in which a portion of the nervous system is destroyed or injured by a nutritional disorder or disorder of metabolism including but not limited to, vitamin B12 deficiency, folic acid deficiency, Wernicke disease, tobacco-alcohol amblyopia, Marchiafava-Bignami disease (primary degeneration of the corpus callosum), and alcoholic cerebellar degeneration;
- (vi) neurological lesions associated with systemic diseases including but not limited to diabetes (diabetic neuropathy, Bell's palsy), systemic lupus crythematosus, carcinoma, or sarcoidosis;
- (vii) lesions caused by toxic substances including alcohol, lead, or particular neurotoxins; and
- (viii) demyelinated lesions in which a portion of the nervous system is destroyed or injured by a demyelinating disease including but not limited to multiple sclerosis, human immunodeficiency virus-associated myelopathy, transverse myelopathy or various etiologies, progressive multifocal leukoencephalopathy, and central pontine myelinolysis.

Therapeutics which are useful according to the invention for treatment of a nervous system disorder may be selected by testing for biological activity in promoting the survival or differentiation of neurons. For example, and not by way of limitation, therapeutics which elicit any of the following effects may be useful according to the invention:

(i) increased survival time of neurons in culture;

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- (ii) increased sprouting of neurons in culture or in vivo;
- (iii) increased production of a neuron-associated molecule in culture or *in vivo*, *e.g.*, choline acetyltransferase or acetylcholinesterase with respect to motor neurons; or
  - (iv) decreased symptoms of neuron dysfunction in vivo.

Such effects may be measured by any method known in the art. In preferred, non-limiting embodiments, increased survival of neurons may be measured by the method set forth in Arakawa et al. (1990, J. Neurosci. 10:3507-3515); increased sprouting of neurons may be detected by methods set forth in Pestronk et al. (1980, Exp. Neurol. 70:65-82) or Brown et al. (1981, Ann. Rev. Neurosci. 4:17-42); increased production of neuron-associated molecules may be measured by bioassay, enzymatic assay, antibody binding, Northern blot assay, etc., depending on the molecule to be measured; and motor neuron dysfunction may be measured by assessing the physical manifestation of motor neuron disorder, e.g., weakness, motor neuron conduction velocity, or functional disability.

In specific embodiments, motor neuron disorders that may be treated according to the invention include but are not limited to disorders such as infarction, infection, exposure to toxin, trauma, surgical damage, degenerative disease or malignancy that may affect motor neurons as well as other components of the nervous system, as well as disorders that selectively affect neurons such as amyotrophic lateral sclerosis, and including but not limited to progressive spinal muscular atrophy, progressive bulbar palsy, primary lateral sclerosis, infantile and juvenile muscular atrophy, progressive bulbar paralysis of childhood (Fazio-Londe syndrome), poliomyelitis and the post polio syndrome, and Hereditary Motorsensory Neuropathy (Charcot-Marie-Tooth Disease).

## 4.10.18 OTHER ACTIVITIES

A polypeptide of the invention may also exhibit one or more of the following additional activities or effects: inhibiting the growth, infection or function of, or killing, infectious agents, including, without limitation, bacteria, viruses, fungi and other parasites; effecting (suppressing or enhancing) bodily characteristics, including, without limitation, height, weight, hair color, eye color, skin, fat to lean ratio or other tissue pigmentation, or organ or body part size or shape (such as, for example, breast augmentation or diminution, change in bone form or shape);

effecting biorhythms or circadian cycles or rhythms; effecting the fertility of male or female subjects; effecting the metabolism, catabolism, anabolism, processing, utilization, storage or elimination of dietary fat, lipid, protein, carbohydrate, vitamins, minerals, co-factors or other nutritional factors or component(s); effecting behavioral characteristics, including, without limitation, appetite, libido, stress, cognition (including cognitive disorders), depression (including depressive disorders) and violent behaviors; providing analgesic effects or other pain reducing effects; promoting differentiation and growth of embryonic stem cells in lineages other than hematopoietic lineages; hormonal or endocrine activity; in the case of enzymes, correcting deficiencies of the enzyme and treating deficiency-related diseases; treatment of hyperproliferative disorders (such as, for example, psoriasis); immunoglobulin-like activity (such as, for example, the ability to bind antigens or complement); and the ability to act as an antigen in a vaccine composition to raise an immune response against such protein or another material or entity which is cross-reactive with such protein.

## 4.10.19 IDENTIFICATION OF POLYMORPHISMS

The demonstration of polymorphisms makes possible the identification of such polymorphisms in human subjects and the pharmacogenetic use of this information for diagnosis and treatment. Such polymorphisms may be associated with, *e.g.*, differential predisposition or susceptibility to various disease states (such as disorders involving inflammation or immune response) or a differential response to drug administration, and this genetic information can be used to tailor preventive or therapeutic treatment appropriately. For example, the existence of a polymorphism associated with a predisposition to inflammation or autoimmune disease makes possible the diagnosis of this condition in humans by identifying the presence of the polymorphism.

Polymorphisms can be identified in a variety of ways known in the art which all generally involve obtaining a sample from a patient, analyzing DNA from the sample, optionally involving isolation or amplification of the DNA, and identifying the presence of the polymorphism in the DNA. For example, PCR may be used to amplify an appropriate fragment of genomic DNA which may then be sequenced. Alternatively, the DNA may be subjected to allele-specific oligonucleotide hybridization (in which appropriate oligonucleotides are hybridized to the DNA under conditions permitting detection of a single base mismatch) or to a single nucleotide extension assay (in which an oligonucleotide that hybridizes immediately adjacent to the position of the polymorphism is extended with one or more labeled nucleotides). In addition, traditional restriction fragment length polymorphism analysis (using restriction enzymes that provide differential digestion of the genomic DNA depending on the presence or

absence of the polymorphism) may be performed. Arrays with nucleotide sequences of the present invention can be used to detect polymorphisms. The array can comprise modified nucleotide sequences of the present invention in order to detect the nucleotide sequences of the present invention. In the alternative, any one of the nucleotide sequences of the present invention can be placed on the array to detect changes from those sequences.

Alternatively a polymorphism resulting in a change in the amino acid sequence could also be detected by detecting a corresponding change in amino acid sequence of the protein, e.g., by an antibody specific to the variant sequence.

## 4.10.20 ARTHRITIS AND INFLAMMATION

The immunosuppressive effects of the compositions of the invention against rheumatoid arthritis is determined in an experimental animal model system. The experimental model system is adjuvant induced arthritis in rats, and the protocol is described by J. Holoshitz, et at., 1983, Science, 219:56, or by B. Waksman et al., 1963, Int. Arch. Allergy Appl. Immunol., 23:129. Induction of the disease can be caused by a single injection, generally intradermally, of a suspension of killed Mycobacterium tuberculosis in complete Freund's adjuvant (CFA). The route of injection can vary, but rats may be injected at the base of the tail with an adjuvant mixture. The polypeptide is administered in phosphate buffered solution (PBS) at a dose of about 1-5 mg/kg. The control consists of administering PBS only.

The procedure for testing the effects of the test compound would consist of intradermally injecting killed Mycobacterium tuberculosis in CFA followed by immediately administering the test compound and subsequent treatment every other day until day 24. At 14, 15, 18, 20, 22, and 24 days after injection of Mycobacterium CFA, an overall arthritis score may be obtained as described by J. Holoskitz above. An analysis of the data would reveal that the test compound would have a dramatic affect on the swelling of the joints as measured by a decrease of the arthritis score.

## 4.11 THERAPEUTIC METHODS

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The compositions (including polypeptide fragments, analogs, variants and antibodies or other binding partners or modulators including antisense polynucleotides) of the invention have numerous applications in a variety of therapeutic methods. Examples of therapeutic applications include, but are not limited to, those exemplified herein.

## **4.11.1 EXAMPLE**

One embodiment of the invention is the administration of an effective amount of the polypeptides or other composition of the invention to individuals affected by a disease or disorder that can be modulated by regulating the peptides of the invention. While the mode of administration is not particularly important, parenteral administration is preferred. An exemplary mode of administration is to deliver an intravenous bolus. The dosage of the polypeptides or other composition of the invention will normally be determined by the prescribing physician. It is to be expected that the dosage will vary according to the age, weight, condition and response of the individual patient. Typically, the amount of polypeptide administered per dose will be in the range of about 0.01µg/kg to 100 mg/kg of body weight, with the preferred dose being about 0.1µg/kg to 10 mg/kg of patient body weight. For parenteral administration, polypeptides of the invention will be formulated in an injectable form combined with a pharmaceutically acceptable parenteral vehicle. Such vehicles are well known in the art and examples include water, saline, Ringer's solution, dextrose solution, and solutions consisting of small amounts of the human serum albumin. The vehicle may contain minor amounts of additives that maintain the isotonicity and stability of the polypeptide or other active ingredient. The preparation of such solutions is within the skill of the art.

# 4.12 PHARMACEUTICAL FORMULATIONS AND ROUTES OF ADMINISTRATION

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A protein or other composition of the present invention (from whatever source derived, including without limitation from recombinant and non-recombinant sources and including antibodies and other binding partners of the polypeptides of the invention) may be administered to a patient in need, by itself, or in pharmaceutical compositions where it is mixed with suitable carriers or excipient(s) at doses to treat or ameliorate a variety of disorders. Such a composition may optionally contain (in addition to protein or other active ingredient and a carrier) diluents, fillers, salts, buffers, stabilizers, solubilizers, and other materials well known in the art. The term "pharmaceutically acceptable" means a non-toxic material that does not interfere with the effectiveness of the biological activity of the active ingredient(s). The characteristics of the carrier will depend on the route of administration. The pharmaceutical composition of the invention may also contain cytokines, lymphokines, or other hematopoietic factors such as M-CSF, GM-CSF, TNF, IL-1, IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, IL-10, IL-11, IL-12, IL-13, IL-14, IL-15, IFN, TNF0, TNF1, TNF2, G-CSF, Meg-CSF, thrombopoietin, stem cell factor, and erythropoietin. In further compositions, proteins of the invention may be combined with other agents beneficial to the treatment of the disease or disorder in question. These agents include various growth factors such as epidermal growth factor (EGF), platelet-derived growth

factor (PDGF), transforming growth factors (TGF- $\alpha$  and TGF- $\beta$ ), insulin-like growth factor (IGF), as well as cytokines described herein.

The pharmaceutical composition may further contain other agents which either enhance the activity of the protein or other active ingredient or complement its activity or use in treatment. Such additional factors and/or agents may be included in the pharmaceutical composition to produce a synergistic effect with protein or other active ingredient of the invention, or to minimize side effects. Conversely, protein or other active ingredient of the present invention may be included in formulations of the particular clotting factor, cytokine, lymphokine, other hematopoietic factor, thrombolytic or anti-thrombotic factor, or anti-inflammatory agent to minimize side effects of the clotting factor, cytokine, lymphokine, other hematopoietic factor, thrombolytic or anti-thrombotic factor, or anti-inflammatory agent (such as IL-1Ra, IL-1 Hy1, IL-1 Hy2, anti-TNF, corticosteroids, immunosuppressive agents). A protein of the present invention may be active in multimers (e.g., heterodimers or homodimers) or complexes with itself or other proteins. As a result, pharmaceutical compositions of the invention may comprise a protein of the invention in such multimeric or complexed form.

As an alternative to being included in a pharmaceutical composition of the invention including a first protein, a second protein or a therapeutic agent may be concurrently administered with the first protein (e.g., at the same time, or at differing times provided that therapeutic concentrations of the combination of agents is achieved at the treatment site). Techniques for formulation and administration of the compounds of the instant application may be found in "Remington's Pharmaceutical Sciences," Mack Publishing Co., Easton, PA, latest edition. A therapeutically effective dose further refers to that amount of the compound sufficient to result in amelioration of symptoms, e.g., treatment, healing, prevention or amelioration of the relevant medical condition, or an increase in rate of treatment, healing, prevention or amelioration of such conditions. When applied to an individual active ingredient, administered alone, a therapeutically effective dose refers to that ingredient alone. When applied to a combination, a therapeutically effective dose refers to combined amounts of the active ingredients that result in the therapeutic effect, whether administered in combination, serially or simultaneously.

In practicing the method of treatment or use of the present invention, a therapeutically effective amount of protein or other active ingredient of the present invention is administered to a mammal having a condition to be treated. Protein or other active ingredient of the present invention may be administered in accordance with the method of the invention either alone or in combination with other therapies such as treatments employing cytokines, lymphokines or other hematopoietic factors. When co- administered with one or more cytokines, lymphokines or other

hematopoietic factors, protein or other active ingredient of the present invention may be administered either simultaneously with the cytokine(s), lymphokine(s), other hematopoietic factor(s), thrombolytic or anti-thrombotic factors, or sequentially. If administered sequentially, the attending physician will decide on the appropriate sequence of administering protein or other active ingredient of the present invention in combination with cytokine(s), lymphokine(s), other hematopoietic factor(s), thrombolytic or anti-thrombotic factors.

#### 4.12.1 ROUTES OF ADMINISTRATION

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Suitable routes of administration may, for example, include oral, rectal, transmucosal, or intestinal administration; parenteral delivery, including intramuscular, subcutaneous, intramedullary injections, as well as intrathecal, direct intraventricular, intravenous, intraperitoneal, intranasal, or intraocular injections. Administration of protein or other active ingredient of the present invention used in the pharmaceutical composition or to practice the method of the present invention can be carried out in a variety of conventional ways, such as oral ingestion, inhalation, topical application or cutaneous, subcutaneous, intraperitoneal, parenteral or intravenous injection. Intravenous administration to the patient is preferred.

Alternately, one may administer the compound in a local rather than systemic manner, for example, via injection of the compound directly into a arthritic joints or in fibrotic tissue, often in a depot or sustained release formulation. In order to prevent the scarring process frequently occurring as complication of glaucoma surgery, the compounds may be administered topically, for example, as eye drops. Furthermore, one may administer the drug in a targeted drug delivery system, for example, in a liposome coated with a specific antibody, targeting, for example, arthritic or fibrotic tissue. The liposomes will be targeted to and taken up selectively by the afflicted tissue.

The polypeptides of the invention are administered by any route that delivers an effective dosage to the desired site of action. The determination of a suitable route of administration and an effective dosage for a particular indication is within the level of skill in the art. Preferably for wound treatment, one administers the therapeutic compound directly to the site. Suitable dosage ranges for the polypeptides of the invention can be extrapolated from these dosages or from similar studies in appropriate animal models. Dosages can then be adjusted as necessary by the clinician to provide maximal therapeutic benefit.

## 4.12.2 COMPOSITIONS/FORMULATIONS

Pharmaceutical compositions for use in accordance with the present invention thus may be formulated in a conventional manner using one or more physiologically acceptable carriers

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comprising excipients and auxiliaries which facilitate processing of the active compounds into preparations which can be used pharmaceutically. These pharmaceutical compositions may be manufactured in a manner that is itself known, e.g., by means of conventional mixing, dissolving, granulating, dragee-making, levigating, emulsifying, encapsulating, entrapping or lyophilizing processes. Proper formulation is dependent upon the route of administration chosen. When a therapeutically effective amount of protein or other active ingredient of the present invention is administered orally, protein or other active ingredient of the present invention will be in the form of a tablet, capsule, powder, solution or elixir. When administered in tablet form, the pharmaceutical composition of the invention may additionally contain a solid carrier such as a gelatin or an adjuvant. The tablet, capsule, and powder contain from about 5 to 95% protein or other active ingredient of the present invention, and preferably from about 25 to 90% protein or other active ingredient of the present invention. When administered in liquid form, a liquid carrier such as water, petroleum, oils of animal or plant origin such as peanut oil, mineral oil, soybean oil, or sesame oil, or synthetic oils may be added. The liquid form of the pharmaceutical composition may further contain physiological saline solution, dextrose or other saccharide solution, or glycols such as ethylene glycol, propylene glycol or polyethylene glycol. When administered in liquid form, the pharmaceutical composition contains from about 0.5 to 90% by weight of protein or other active ingredient of the present invention, and preferably from about 1 to 50% protein or other active ingredient of the present invention.

When a therapeutically effective amount of protein or other active ingredient of the present invention is administered by intravenous, cutaneous or subcutaneous injection, protein or other active ingredient of the present invention will be in the form of a pyrogen-free, parenterally acceptable aqueous solution. The preparation of such parenterally acceptable protein or other active ingredient solutions, having due regard to pH, isotonicity, stability, and the like, is within the skill in the art. A preferred pharmaceutical composition for intravenous, cutaneous, or subcutaneous injection should contain, in addition to protein or other active ingredient of the present invention, an isotonic vehicle such as Sodium Chloride Injection, Ringer's Injection, Dextrose Injection, Dextrose and Sodium Chloride Injection, Lactated Ringer's Injection, or other vehicle as known in the art. The pharmaceutical composition of the present invention may also contain stabilizers, preservatives, buffers, antioxidants, or other additives known to those of skill in the art. For injection, the agents of the invention may be formulated in aqueous solutions, preferably in physiologically compatible buffers such as Hanks's solution, Ringer's solution, or physiological saline buffer. For transmucosal administration, penetrants appropriate to the barrier to be permeated are used in the formulation. Such penetrants are generally known in the art.

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For oral administration, the compounds can be formulated readily by combining the active compounds with pharmaceutically acceptable carriers well known in the art. Such carriers enable the compounds of the invention to be formulated as tablets, pills, dragees, capsules, liquids, gels, syrups, slurries, suspensions and the like, for oral ingestion by a patient to be treated. Pharmaceutical preparations for oral use can be obtained from a solid excipient, optionally grinding a resulting mixture, and processing the mixture of granules, after adding suitable auxiliaries, if desired, to obtain tablets or dragee cores. Suitable excipients are, in particular, fillers such as sugars, including lactose, sucrose, mannitol, or sorbitol; cellulose preparations such as, for example, maize starch, wheat starch, rice starch, potato starch, gelatin, gum tragacanth, methyl cellulose, hydroxypropylmethyl-cellulose, sodium carboxymethylcellulose, and/or polyvinylpyrrolidone (PVP). If desired, disintegrating agents may be added, such as the cross-linked polyvinyl pyrrolidone, agar, or alginic acid or a salt thereof such as sodium alginate. Dragee cores are provided with suitable coatings. For this purpose, concentrated sugar solutions may be used, which may optionally contain gum arabic, tale, polyvinyl pyrrolidone, carbopol gel, polyethylene glycol, and/or titanium dioxide, lacquer solutions, and suitable organic solvents or solvent mixtures. Dyestuffs or pigments may be added to the tablets or dragee coatings for identification or to characterize different combinations of active compound doses.

Pharmaceutical preparations which can be used orally include push-fit capsules made of gelatin, as well as soft, sealed capsules made of gelatin and a plasticizer, such as glycerol or sorbitol. The push-fit capsules can contain the active ingredients in admixture with filler such as lactose, binders such as starches, and/or lubricants such as talc or magnesium stearate and, optionally, stabilizers. In soft capsules, the active compounds may be dissolved or suspended in suitable liquids, such as fatty oils, liquid paraffin, or liquid polyethylene glycols. In addition, stabilizers may be added. All formulations for oral administration should be in dosages suitable for such administration. For buccal administration, the compositions may take the form of tablets or lozenges formulated in conventional manner.

For administration by inhalation, the compounds for use according to the present invention are conveniently delivered in the form of an aerosol spray presentation from pressurized packs or a nebuliser, with the use of a suitable propellant, *e.g.*, dichlorodifluoromethane, trichlorofluoromethane, dichlorotetrafluoroethane, carbon dioxide or other suitable gas. In the case of a pressurized aerosol the dosage unit may be determined by providing a valve to deliver a metered amount. Capsules and cartridges of, *e.g.*, gelatin for use in an inhaler or insufflator may be formulated containing a powder mix of the compound and a suitable powder base such as lactose or starch. The compounds may be formulated for parenteral

administration by injection, *e.g.*, by bolus injection or continuous infusion. Formulations for injection may be presented in unit dosage form, *e.g.*, in ampules or in multi-dose containers, with an added preservative. The compositions may take such forms as suspensions, solutions or emulsions in oily or aqueous vehicles, and may contain formulatory agents such as suspending, stabilizing and/or dispersing agents.

Pharmaceutical formulations for parenteral administration include aqueous solutions of the active compounds in water-soluble form. Additionally, suspensions of the active compounds may be prepared as appropriate oily injection suspensions. Suitable lipophilic solvents or vehicles include fatty oils such as sesame oil, or synthetic fatty acid esters, such as ethyl oleate or triglycerides, or liposomes. Aqueous injection suspensions may contain substances which increase the viscosity of the suspension, such as sodium carboxymethyl cellulose, sorbitol, or dextran. Optionally, the suspension may also contain suitable stabilizers or agents which increase the solubility of the compounds to allow for the preparation of highly concentrated solutions. Alternatively, the active ingredient may be in powder form for constitution with a suitable vehicle, *e.g.*, sterile pyrogen-free water, before use.

The compounds may also be formulated in rectal compositions such as suppositories or retention enemas, *e.g.*, containing conventional suppository bases such as cocoa butter or other glycerides. In addition to the formulations described previously, the compounds may also be formulated as a depot preparation. Such long acting formulations may be administered by implantation (for example subcutaneously or intramuscularly) or by intramuscular injection. Thus, for example, the compounds may be formulated with suitable polymeric or hydrophobic materials (for example as an emulsion in an acceptable oil) or ion exchange resins, or as sparingly soluble derivatives, for example, as a sparingly soluble salt.

A pharmaceutical carrier for the hydrophobic compounds of the invention is a co-solvent system comprising benzyl alcohol, a nonpolar surfactant, a water-miscible organic polymer, and an aqueous phase. The co-solvent system may be the VPD co-solvent system. VPD is a solution of 3% w/v benzyl alcohol, 8% w/v of the nonpolar surfactant polysorbate 80, and 65% w/v polyethylene glycol 300, made up to volume in absolute ethanol. The VPD co-solvent system (VPD:5W) consists of VPD diluted 1:1 with a 5% dextrose in water solution. This co-solvent system dissolves hydrophobic compounds well, and itself produces low toxicity upon systemic administration. Naturally, the proportions of a co-solvent system may be varied considerably without destroying its solubility and toxicity characteristics. Furthermore, the identity of the co-solvent components may be varied: for example, other low-toxicity nonpolar surfactants may be used instead of polysorbate 80; the fraction size of polyethylene glycol may be varied; other biocompatible polymers may replace polyethylene glycol, e.g. polyvinyl pyrrolidone; and other

sugars or polysaccharides may substitute for dextrose. Alternatively, other delivery systems for hydrophobic pharmaceutical compounds may be employed. Liposomes and emulsions are well known examples of delivery vehicles or carriers for hydrophobic drugs. Certain organic solvents such as dimethylsulfoxide also may be employed, although usually at the cost of greater toxicity.

Additionally, the compounds may be delivered using a sustained-release system, such as semipermeable matrices of solid hydrophobic polymers containing the therapeutic agent. Various types of sustained-release materials have been established and are well known by those skilled in the art. Sustained-release capsules may, depending on their chemical nature, release the compounds for a few weeks up to over 100 days. Depending on the chemical nature and the biological stability of the therapeutic reagent, additional strategies for protein or other active ingredient stabilization may be employed.

The pharmaceutical compositions also may comprise suitable solid or gel phase carriers or excipients. Examples of such carriers or excipients include but are not limited to calcium carbonate, calcium phosphate, various sugars, starches, cellulose derivatives, gelatin, and polymers such as polyethylene glycols. Many of the active ingredients of the invention may be provided as salts with pharmaceutically compatible counter ions. Such pharmaceutically acceptable base addition salts are those salts which retain the biological effectiveness and properties of the free acids and which are obtained by reaction with inorganic or organic bases such as sodium hydroxide, magnesium hydroxide, ammonia, trialkylamine, dialkylamine, monoalkylamine, dibasic amino acids, sodium acetate, potassium benzoate, triethanol amine and the like.

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The pharmaceutical composition of the invention may be in the form of a complex of the protein(s) or other active ingredient(s) of present invention along with protein or peptide antigens. The protein and/or peptide antigen will deliver a stimulatory signal to both B and T lymphocytes. B lymphocytes will respond to antigen through their surface immunoglobulin receptor. T lymphocytes will respond to antigen through the T cell receptor (TCR) following presentation of the antigen by MHC proteins. MHC and structurally related proteins including those encoded by class I and class II MHC genes on host cells will serve to present the peptide antigen(s) to T lymphocytes. The antigen components could also be supplied as purified MHC-peptide complexes alone or with co-stimulatory molecules that can directly signal T cells. Alternatively antibodies able to bind surface immunoglobulin and other molecules on B cells as well as antibodies able to bind the TCR and other molecules on T cells can be combined with the pharmaceutical composition of the invention.

The pharmaceutical composition of the invention may be in the form of a liposome in which protein of the present invention is combined, in addition to other pharmaceutically

acceptable carriers, with amphipathic agents such as lipids which exist in aggregated form as micelles, insoluble monolayers, liquid crystals, or lamellar layers in aqueous solution. Suitable lipids for liposomal formulation include, without limitation, monoglycerides, diglycerides, sulfatides, lysolecithins, phospholipids, saponin, bile acids, and the like. Preparation of such liposomal formulations is within the level of skill in the art, as disclosed, for example, in U.S. Patent Nos. 4,235,871; 4,501,728; 4,837,028; and 4,737,323, all of which are incorporated herein by reference.

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The amount of protein or other active ingredient of the present invention in the pharmaceutical composition of the present invention will depend upon the nature and severity of the condition being treated, and on the nature of prior treatments which the patient has undergone. Ultimately, the attending physician will decide the amount of protein or other active ingredient of the present invention with which to treat each individual patient. Initially, the attending physician will administer low doses of protein or other active ingredient of the present invention and observe the patient's response. Larger doses of protein or other active ingredient of the present invention may be administered until the optimal therapeutic effect is obtained for the patient, and at that point the dosage is not increased further. It is contemplated that the various pharmaceutical compositions used to practice the method of the present invention should contain about 0.01 µg to about 100 mg (preferably about 0.1 µg to about 10 mg, more preferably about 0.1 µg to about 1 mg) of protein or other active ingredient of the present invention per kg body weight. For compositions of the present invention which are useful for bone, cartilage, tendon or ligament regeneration, the therapeutic method includes administering the composition topically, systematically, or locally as an implant or device. When administered, the therapeutic composition for use in this invention is, of course, in a pyrogen-free, physiologically acceptable form. Further, the composition may desirably be encapsulated or injected in a viscous form for delivery to the site of bone, cartilage or tissue damage. Topical administration may be suitable for wound healing and tissue repair. Therapeutically useful agents other than a protein or other active ingredient of the invention which may also optionally be included in the composition as described above, may alternatively or additionally, be administered simultaneously or sequentially with the composition in the methods of the invention. Preferably for bone and/or cartilage formation, the composition would include a matrix capable of delivering the protein-containing or other active ingredient-containing composition to the site of bone and/or cartilage damage, providing a structure for the developing bone and cartilage and optimally capable of being resorbed into the body. Such matrices may be formed of materials presently in use for other implanted medical applications.

The choice of matrix material is based on biocompatibility, biodegradability, mechanical properties, cosmetic appearance and interface properties. The particular application of the compositions will define the appropriate formulation. Potential matrices for the compositions may be biodegradable and chemically defined calcium sulfate, tricalcium phosphate, hydroxyapatite, polylactic acid, polyglycolic acid and polyanhydrides. Other potential materials are biodegradable and biologically well-defined, such as bone or dermal collagen. Further matrices are comprised of pure proteins or extracellular matrix components. Other potential matrices are nonbiodegradable and chemically defined, such as sintered hydroxyapatite, bioglass, aluminates, or other ceramics. Matrices may be comprised of combinations of any of the above 10 mentioned types of material, such as polylactic acid and hydroxyapatite or collagen and tricalcium phosphate. The bioceramics may be altered in composition, such as in calcium-aluminate-phosphate and processing to alter pore size, particle size, particle shape, and biodegradability. Presently preferred is a 50:50 (mole weight) copolymer of lactic acid and glycolic acid in the form of porous particles having diameters ranging from 150 to 800 microns. 15 In some applications, it will be useful to utilize a sequestering agent, such as carboxymethyl cellulose or autologous blood clot, to prevent the protein compositions from disassociating from the matrix.

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A preferred family of sequestering agents is cellulosic materials such as alkylcelluloses (including hydroxyalkylcelluloses), including methylcellulose, ethylcellulose, hydroxyethylcellulose, hydroxypropylcellulose, hydroxypropyl-methylcellulose, and 20 carboxymethylcellulose, the most preferred being cationic salts of carboxymethylcellulose (CMC). Other preferred sequestering agents include hyaluronic acid, sodium alginate, poly(ethylene glycol), polyoxyethylene oxide, carboxyvinyl polymer and poly(vinyl alcohol). The amount of sequestering agent useful herein is 0.5-20 wt %, preferably 1-10 wt % based on 25 total formulation weight, which represents the amount necessary to prevent desorption of the protein from the polymer matrix and to provide appropriate handling of the composition, yet not so much that the progenitor cells are prevented from infiltrating the matrix, thereby providing the protein the opportunity to assist the osteogenic activity of the progenitor cells. In further compositions, proteins or other active ingredients of the invention may be combined with other 30 agents beneficial to the treatment of the bone and/or cartilage defect, wound, or tissue in question. These agents include various growth factors such as epidermal growth factor (EGF), platelet derived growth factor (PDGF), transforming growth factors (TGF-α and TGF-β), and insulin-like growth factor (IGF).

The therapeutic compositions are also presently valuable for veterinary applications. Particularly domestic animals and thoroughbred horses, in addition to humans, are desired

patients for such treatment with proteins or other active ingredients of the present invention. The dosage regimen of a protein-containing pharmaceutical composition to be used in tissue regeneration will be determined by the attending physician considering various factors which modify the action of the proteins, *e.g.*, amount of tissue weight desired to be formed, the site of damage, the condition of the damaged tissue, the size of a wound, type of damaged tissue (*e.g.*, bone), the patient's age, sex, and diet, the severity of any infection, time of administration and other clinical factors. The dosage may vary with the type of matrix used in the reconstitution and with inclusion of other proteins in the pharmaceutical composition. For example, the addition of other known growth factors, such as IGF I (insulin like growth factor I), to the final composition, may also effect the dosage. Progress can be monitored by periodic assessment of tissue/bone growth and/or repair, for example, X-rays, histomorphometric determinations and tetracycline labeling.

Polynucleotides of the present invention can also be used for gene therapy. Such polynucleotides can be introduced either in vivo or ex vivo into cells for expression in a mammalian subject. Polynucleotides of the invention may also be administered by other known methods for introduction of nucleic acid into a cell or organism (including, without limitation, in the form of viral vectors or naked DNA). Cells may also be cultured ex vivo in the presence of proteins of the present invention in order to proliferate or to produce a desired effect on or activity in such cells. Treated cells can then be introduced in vivo for therapeutic purposes.

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## 4.12.3 EFFECTIVE DOSAGE

Pharmaceutical compositions suitable for use in the present invention include compositions wherein the active ingredients are contained in an effective amount to achieve its intended purpose. More specifically, a therapeutically effective amount means an amount effective to prevent development of or to alleviate the existing symptoms of the subject being treated. Determination of the effective amount is well within the capability of those skilled in the art, especially in light of the detailed disclosure provided herein. For any compound used in the method of the invention, the therapeutically effective dose can be estimated initially from appropriate in vitro assays. For example, a dose can be formulated in animal models to achieve a circulating concentration range that can be used to more accurately determine useful doses in humans. For example, a dose can be formulated in animal models to achieve a circulating concentration range that includes the IC<sub>50</sub> as determined in cell culture (*i.e.*, the concentration of the test compound which achieves a half-maximal inhibition of the protein's biological activity). Such information can be used to more accurately determine useful doses in humans.

A therapeutically effective dose refers to that amount of the compound that results in amelioration of symptoms or a prolongation of survival in a patient. Toxicity and therapeutic efficacy of such compounds can be determined by standard pharmaceutical procedures in cell cultures or experimental animals, e.g., for determining the LD<sub>50</sub> (the dose lethal to 50% of the population) and the ED<sub>50</sub> (the dose therapeutically effective in 50% of the population). The dose ratio between toxic and therapeutic effects is the therapeutic index and it can be expressed as the ratio between LD<sub>50</sub> and ED<sub>50</sub>. Compounds which exhibit high therapeutic indices are preferred. The data obtained from these cell culture assays and animal studies can be used in formulating a range of dosage for use in human. The dosage of such compounds lies preferably within a range of circulating concentrations that include the  $ED_{50}$  with little or no toxicity. The dosage may vary within this range depending upon the dosage form employed and the route of administration utilized. The exact formulation, route of administration and dosage can be chosen by the individual physician in view of the patient's condition. See, e.g., Fingl et al., 1975, in "The Pharmacological Basis of Therapeutics", Ch. 1 p.1. Dosage amount and interval may be adjusted individually to provide plasma levels of the active moiety which are sufficient to maintain the desired effects, or minimal effective concentration (MEC). The MEC will vary for each compound but can be estimated from in vitro data. Dosages necessary to achieve the MEC will depend on individual characteristics and route of administration. However, HPLC assays or bioassays can be used to determine plasma concentrations.

Dosage intervals can also be determined using MEC value. Compounds should be administered using a regimen which maintains plasma levels above the MEC for 10-90% of the time, preferably between 30-90% and most preferably between 50-90%. In cases of local administration or selective uptake, the effective local concentration of the drug may not be related to plasma concentration.

An exemplary dosage regimen for polypeptides or other compositions of the invention will be in the range of about 0.01  $\mu$ g/kg to 100 mg/kg of body weight daily, with the preferred dose being about 0.1  $\mu$ g/kg to 25 mg/kg of patient body weight daily, varying in adults and children. Dosing may be once daily, or equivalent doses may be delivered at longer or shorter intervals.

The amount of composition administered will, of course, be dependent on the subject being treated, on the subject's age and weight, the severity of the affliction, the manner of administration and the judgment of the prescribing physician.

#### 4.12.4 PACKAGING

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The compositions may, if desired, be presented in a pack or dispenser device which may contain one or more unit dosage forms containing the active ingredient. The pack may, for example, comprise metal or plastic foil, such as a blister pack. The pack or dispenser device may be accompanied by instructions for administration. Compositions comprising a compound of the invention formulated in a compatible pharmaceutical carrier may also be prepared, placed in an appropriate container, and labeled for treatment of an indicated condition.

#### 4.13 ANTIBODIES

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Also included in the invention are antibodies to proteins, or fragments of proteins of the invention. The term "antibody" as used herein refers to immunoglobulin molecules and immunologically active portions of immunoglobulin (Ig) molecules, *i.e.*, molecules that contain an antigen binding site that specifically binds (immunoreacts with) an antigen. Such antibodies include, but are not limited to, polyclonal, monoclonal, chimeric, single chain,  $F_{ab}$ ,  $F_{ab}$  and  $F_{(ab)/2}$  fragments, and an  $F_{ab}$  expression library. In general, an antibody molecule obtained from humans relates to any of the classes IgG, IgM, IgA, IgE and IgD, which differ from one another by the nature of the heavy chain present in the molecule. Certain classes have subclasses as well, such as  $IgG_1$ ,  $IgG_2$ , and others. Furthermore, in humans, the light chain may be a kappa chain or a lambda chain. Reference herein to antibodies includes a reference to all such classes, subclasses and types of human antibody species.

An isolated related protein of the invention may be intended to serve as an antigen, or a portion or fragment thereof, and additionally can be used as an immunogen to generate antibodies that immunospecifically bind the antigen, using standard techniques for polyclonal and monoclonal antibody preparation. The full-length protein can be used or, alternatively, the invention provides antigenic peptide fragments of the antigen for use as immunogens. An antigenic peptide fragment comprises at least 6 amino acid residues of the amino acid sequence of the full length protein, (for example the amino acid sequence shown in SEQ ID NO: 1351), and encompasses an epitope thereof such that an antibody raised against the peptide forms a specific immune complex with the full length protein or with any fragment that contains the epitope. Preferably, the antigenic peptide comprises at least 10 amino acid residues, or at least 15 amino acid residues, or at least 20 amino acid residues. Preferred epitopes encompassed by the antigenic peptide are regions of the protein that are located on its surface; commonly these are hydrophilic regions.

In certain embodiments of the invention, at least one epitope encompassed by the antigenic peptide is a region of -related protein that is located on the surface of the protein, e.g., a hydrophilic region. A hydrophobicity analysis of the human related protein sequence will

indicate which regions of a related protein are particularly hydrophilic and, therefore, are likely to encode surface residues useful for targeting antibody production. As a means for targeting antibody production, hydropathy plots showing regions of hydrophilicity and hydrophobicity may be generated by any method well known in the art, including, for example, the Kyte Doolittle or the Hopp Woods methods, either with or without Fourier transformation. See, e.g., Hopp and Woods, 1981, Proc. Nat. Acad. Sci. USA 78: 3824-3828; Kyte and Doolittle 1982, J. Mol. Biol. 157: 105-142, each of which is incorporated herein by reference in its entirety. Antibodies that are specific for one or more domains within an antigenic protein, or derivatives, fragments, analogs or homologs thereof, are also provided herein.

A protein of the invention, or a derivative, fragment, analog, homolog or ortholog thereof, may be utilized as an immunogen in the generation of antibodies that immunospecifically bind these protein components.

Various procedures known within the art may be used for the production of polyclonal or monoclonal antibodies directed against a protein of the invention, or against derivatives, fragments, analogs homologs or orthologs thereof (see, for example, Antibodies: A Laboratory Manual, Harlow E, and Lane D, 1988, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, incorporated herein by reference). Some of these antibodies are discussed below.

## 5.13.1 Polyclonal Antibodies

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For the production of polyclonal antibodies, various suitable host animals (e.g., rabbit, goat, mouse or other mammal) may be immunized by one or more injections with the native protein, a synthetic variant thereof, or a derivative of the foregoing. An appropriate immunogenic preparation can contain, for example, the naturally occurring immunogenic protein, a chemically synthesized polypeptide representing the immunogenic protein, or a recombinantly expressed immunogenic protein. Furthermore, the protein may be conjugated to a second protein known to be immunogenic in the mammal being immunized. Examples of such immunogenic proteins include but are not limited to keyhole limpet hemocyanin, serum albumin, bovine thyroglobulin, and soybean trypsin inhibitor. The preparation can further include an adjuvant. Various adjuvants used to increase the immunological response include, but are not limited to, Freund's (complete and incomplete), mineral gels (e.g., aluminum hydroxide), surface active substances (e.g., lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, dinitrophenol, etc.), adjuvants usable in humans such as Bacille Calmette-Guerin and Corynebacterium parvum, or similar immunostimulatory agents. Additional examples of adjuvants which can be employed include MPL-TDM adjuvant (monophosphoryl Lipid A, 35 synthetic trehalose dicorynomycolate).

The polyclonal antibody molecules directed against the immunogenic protein can be isolated from the mammal (e.g., from the blood) and further purified by well known techniques, such as affinity chromatography using protein A or protein G, which provide primarily the IgG fraction of immune serum. Subsequently, or alternatively, the specific antigen which is the target of the immunoglobulin sought, or an epitope thereof, may be immobilized on a column to purify the immune specific antibody by immunoaffinity chromatography. Purification of immunoglobulins is discussed, for example, by D. Wilkinson (The Scientist, published by The Scientist, Inc., Philadelphia PA, Vol. 14, No. 8 (April 17, 2000), pp. 25-28).

#### 5.13.2 Monoclonal Antibodies

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The term "monoclonal antibody" (MAb) or "monoclonal antibody composition", as used herein, refers to a population of antibody molecules that contain only one molecular species of antibody molecule consisting of a unique light chain gene product and a unique heavy chain gene product. In particular, the complementarity determining regions (CDRs) of the monoclonal antibody are identical in all the molecules of the population. MAbs thus contain an antigen binding site capable of immunoreacting with a particular epitope of the antigen characterized by a unique binding affinity for it.

Monoclonal antibodies can be prepared using hybridoma methods, such as those described by Kohler and Milstein, <u>Nature</u>, <u>256</u>:495 (1975). In a hybridoma method, a mouse, hamster, or other appropriate host animal, is typically immunized with an immunizing agent to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the immunizing agent. Alternatively, the lymphocytes can be immunized in vitro.

The immunizing agent will typically include the protein antigen, a fragment thereof or a fusion protein thereof. Generally, either peripheral blood lymphocytes are used if cells of human origin are desired, or spleen cells or lymph node cells are used if non-human mammalian sources are desired. The lymphocytes are then fused with an immortalized cell line using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell (Goding, Monoclonal Antibodies: Principles and Practice, Academic Press, (1986) pp. 59-103). Immortalized cell lines are usually transformed mammalian cells, particularly myeloma cells of rodent, bovine and human origin. Usually, rat or mouse myeloma cell lines are employed. The hybridoma cells can be cultured in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, immortalized cells. For example, if the parental cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine ("HAT medium"), which substances prevent the growth of HGPRT-deficient cells.

Preferred immortalized cell lines are those that fuse efficiently, support stable high level expression of antibody by the selected antibody-producing cells, and are sensitive to a medium such as HAT medium. More preferred immortalized cell lines are murine myeloma lines, which can be obtained, for instance, from the Salk Institute Cell Distribution Center, San Diego, California and the American Type Culture Collection, Manassas, Virginia. Human myeloma and mouse-human heteromyeloma cell lines also have been described for the production of human monoclonal antibodies (Kozbor, <u>J. Immunol., 133</u>:3001 (1984); Brodeur et al., <u>Monoclonal Antibody Production Techniques and Applications</u>, Marcel Dekker, Inc., New York, (1987) pp. 51-63).

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The culture medium in which the hybridoma cells are cultured can then be assayed for the presence of monoclonal antibodies directed against the antigen. Preferably, the binding specificity of monoclonal antibodies produced by the hybridoma cells is determined by immunoprecipitation or by an in vitro binding assay, such as radioimmunoassay (RIA) or enzyme-linked immunoabsorbent assay (ELISA). Such techniques and assays are known in the art. The binding affinity of the monoclonal antibody can, for example, be determined by the Scatchard analysis of Munson and Pollard, <u>Anal. Biochem.</u>, <u>107</u>:220 (1980). Preferably, antibodies having a high degree of specificity and a high binding affinity for the target antigen are isolated.

After the desired hybridoma cells are identified, the clones can be subcloned by limiting dilution procedures and grown by standard methods. Suitable culture media for this purpose include, for example, Dulbecco's Modified Eagle's Medium and RPMI-1640 medium. Alternatively, the hybridoma cells can be grown in vivo as ascites in a mammal.

The monoclonal antibodies secreted by the subclones can be isolated or purified from the culture medium or ascites fluid by conventional immunoglobulin purification procedures such as, for example, protein A-Sepharose, hydroxylapatite chromatography, gel electrophoresis, dialysis, or affinity chromatography.

The monoclonal antibodies can also be made by recombinant DNA methods, such as those described in U.S. Patent No. 4,816,567. DNA encoding the monoclonal antibodies of the invention can be readily isolated and sequenced using conventional procedures (e.g., by using oligonucleotide probes that are capable of binding specifically to genes encoding the heavy and light chains of murine antibodies). The hybridoma cells of the invention serve as a preferred source of such DNA. Once isolated, the DNA can be placed into expression vectors, which are then transfected into host cells such as simian COS cells, Chinese hamster ovary (CHO) cells, or myeloma cells that do not otherwise produce immunoglobulin protein, to obtain the synthesis of monoclonal antibodies in the recombinant host cells. The DNA also can be modified, for

example, by substituting the coding sequence for human heavy and light chain constant domains in place of the homologous murine sequences (U.S. Patent No. 4,816,567; Morrison, Nature 368, 812-13 (1994)) or by covalently joining to the immunoglobulin coding sequence all or part of the coding sequence for a non-immunoglobulin polypeptide. Such a non-immunoglobulin polypeptide can be substituted for the constant domains of an antibody of the invention, or can be substituted for the variable domains of one antigen-combining site of an antibody of the invention to create a chimeric bivalent antibody.

## 5.13.2 Humanized Antibodies

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The antibodies directed against the protein antigens of the invention can further comprise humanized antibodies or human antibodies. These antibodies are suitable for administration to humans without engendering an immune response by the human against the administered immunoglobulin. Humanized forms of antibodies are chimeric immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')2 or other antigenbinding subsequences of antibodies) that are principally comprised of the sequence of a human immunoglobulin, and contain minimal sequence derived from a non-human immunoglobulin. Humanization can be performed following the method of Winter and co-workers (Jones et al., Nature, 321:522-525 (1986); Riechmann et al., Nature, 332:323-327 (1988); Verhoeyen et al., Science, 239:1534-1536 (1988)), by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. (See also U.S. Patent No. 5,225,539.) In some instances, Fy framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Humanized antibodies can also comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the framework regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin (Jones et al., 1986; Riechmann et al., 1988; and Presta, Curr. Op. Struct. Biol., 2:593-596 (1992)).

#### 5.13.3 Human Antibodies

Fully human antibodies relate to antibody molecules in which essentially the entire sequences of both the light chain and the heavy chain, including the CDRs, arise from human genes. Such antibodies are termed "human antibodies", or "fully human antibodies" herein.

Human monoclonal antibodies can be prepared by the trioma technique; the human B-cell hybridoma technique (see Kozbor, et al., 1983 Immunol Today 4: 72) and the EBV hybridoma technique to produce human monoclonal antibodies (see Cole, et al., 1985 In: Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc., pp. 77-96). Human monoclonal antibodies may be utilized in the practice of the present invention and may be produced by using human hybridomas (see Cote, et al., 1983. Proc Natl Acad Sci USA 80: 2026-2030) or by transforming human B-cells with Epstein Barr Virus in vitro (see Cole, et al., 1985 In: Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc., pp. 77-96).

In addition, human antibodies can also be produced using additional techniques, including phage display libraries (Hoogenboom and Winter, J. Mol. Biol., 227:381 (1991); Marks et al., J. Mol. Biol., 222:581 (1991)). Similarly, human antibodies can be made by introducing human immunoglobulin loci into transgenic animals, e.g., mice in which the endogenous immunoglobulin genes have been partially or completely inactivated. Upon challenge, human antibody production is observed, which closely resembles that seen in humans in all respects, including gene rearrangement, assembly, and antibody repertoire. This approach is described, for example, in U.S. Patent Nos. 5,545,807; 5,545,806; 5,569,825; 5,625,126; 5,633,425; 5,661,016, and in Marks et al. (Bio/Technology 10, 779-783 (1992)); Lonberg et al. (Nature 368 856-859 (1994)); Morrison (Nature 368, 812-13 (1994)); Fishwild et al., (Nature Biotechnology 14, 845-51 (1996)); Neuberger (Nature Biotechnology 14, 826 (1996)); and Lonberg and Huszar (Intern. Rev. Immunol. 13 65-93 (1995)).

Human antibodies may additionally be produced using transgenic nonhuman animals which are modified so as to produce fully human antibodies rather than the animal's endogenous antibodies in response to challenge by an antigen. (See PCT publication WO94/02602). The endogenous genes encoding the heavy and light immunoglobulin chains in the nonhuman host have been incapacitated, and active loci encoding human heavy and light chain immunoglobulins are inserted into the host's genome. The human genes are incorporated, for example, using yeast artificial chromosomes containing the requisite human DNA segments. An animal which provides all the desired modifications is then obtained as progeny by crossbreeding intermediate transgenic animals containing fewer than the full complement of the modifications. The preferred embodiment of such a nonhuman animal is a mouse, and is termed the Xenomouse<sup>TM</sup> as disclosed in PCT publications WO 96/33735 and WO 96/34096. This animal produces B cells which secrete fully human immunoglobulins. The antibodies can be obtained directly from the animal after immunization with an immunogen of interest, as, for example, a preparation of a polyclonal antibody, or alternatively from immortalized B cells derived from the animal, such as hybridomas producing monoclonal antibodies. Additionally, the genes encoding the

immunoglobulins with human variable regions can be recovered and expressed to obtain the antibodies directly, or can be further modified to obtain analogs of antibodies such as, for example, single chain Fv molecules.

An example of a method of producing a nonhuman host, exemplified as a mouse, lacking expression of an endogenous immunoglobulin heavy chain is disclosed in U.S. Patent No. 5,939,598. It can be obtained by a method including deleting the J segment genes from at least one endogenous heavy chain locus in an embryonic stem cell to prevent rearrangement of the locus and to prevent formation of a transcript of a rearranged immunoglobulin heavy chain locus, the deletion being effected by a targeting vector containing a gene encoding a selectable marker; and producing from the embryonic stem cell a transgenic mouse whose somatic and germ cells contain the gene encoding the selectable marker.

A method for producing an antibody of interest, such as a human antibody, is disclosed in U.S. Patent No. 5,916,771. It includes introducing an expression vector that contains a nucleotide sequence encoding a heavy chain into one mammalian host cell in culture, introducing an expression vector containing a nucleotide sequence encoding a light chain into another mammalian host cell, and fusing the two cells to form a hybrid cell. The hybrid cell expresses an antibody containing the heavy chain and the light chain.

In a further improvement on this procedure, a method for identifying a clinically relevant epitope on an immunogen, and a correlative method for selecting an antibody that binds immunospecifically to the relevant epitope with high affinity, are disclosed in PCT publication WO 99/53049.

## 5.13.4 Fab Fragment's and Single Chain Antibodies

According to the invention, techniques can be adapted for the production of single-chain antibodies specific to an antigenic protein of the invention (see *e.g.*, U.S. Patent No. 4,946,778). In addition, methods can be adapted for the construction of  $F_{ab}$  expression libraries (see *e.g.*, Huse, et al., 1989 Science 246: 1275-1281) to allow rapid and effective identification of monoclonal  $F_{ab}$  fragments with the desired specificity for a protein or derivatives, fragments, analogs or homologs thereof. Antibody fragments that contain the idiotypes to a protein antigen may be produced by techniques known in the art including, but not limited to: (i) an  $F_{(ab')2}$  fragment produced by pepsin digestion of an antibody molecule; (ii) an  $F_{ab}$  fragment generated by reducing the disulfide bridges of an  $F_{(ab')2}$  fragment; (iii) an  $F_{ab}$  fragment generated by the treatment of the antibody molecule with papain and a reducing agent and (iv)  $F_v$  fragments.

## 5.13.5 Bispecific Antibodies

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Bispecific antibodies are monoclonal, preferably human or humanized, antibodies that have binding specificities for at least two different antigens. In the present case, one of the binding specificities is for an antigenic protein of the invention. The second binding target is any other antigen, and advantageously is a cell-surface protein or receptor or receptor subunit.

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Methods for making bispecific antibodies are known in the art. Traditionally, the recombinant production of bispecific antibodies is based on the co-expression of two immunoglobulin heavy-chain/light-chain pairs, where the two heavy chains have different specificities (Milstein and Cuello, Nature, 305:537-539 (1983)). Because of the random assortment of immunoglobulin heavy and light chains, these hybridomas (quadromas) produce a potential mixture of ten different antibody molecules, of which only one has the correct bispecific structure. The purification of the correct molecule is usually accomplished by affinity chromatography steps. Similar procedures are disclosed in WO 93/08829, published 13 May 1993, and in Traunecker *et al.*, 1991 *EMBO J.*, 10:3655-3659.

Antibody variable domains with the desired binding specificities (antibody-antigen combining sites) can be fused to immunoglobulin constant domain sequences. The fusion preferably is with an immunoglobulin heavy-chain constant domain, comprising at least part of the hinge, CH2, and CH3 regions. It is preferred to have the first heavy-chain constant region (CH1) containing the site necessary for light-chain binding present in at least one of the fusions. DNAs encoding the immunoglobulin heavy-chain fusions and, if desired, the immunoglobulin light chain, are inserted into separate expression vectors, and are co-transfected into a suitable host organism. For further details of generating bispecific antibodies see, for example, Suresh et al., Methods in Enzymology, 121:210 (1986).

According to another approach described in WO 96/27011, the interface between a pair of antibody molecules can be engineered to maximize the percentage of heterodimers which are recovered from recombinant cell culture. The preferred interface comprises at least a part of the CH3 region of an antibody constant domain. In this method, one or more small amino acid side chains from the interface of the first antibody molecule are replaced with larger side chains (e.g. tyrosine or tryptophan). Compensatory "cavities" of identical or similar size to the large side chain(s) are created on the interface of the second antibody molecule by replacing large amino acid side chains with smaller ones (e.g. alanine or threonine). This provides a mechanism for increasing the yield of the heterodimer over other unwanted end-products such as homodimers.

Bispecific antibodies can be prepared as full length antibodies or antibody fragments (e.g. F(ab')<sub>2</sub> bispecific antibodies). Techniques for generating bispecific antibodies from antibody fragments have been described in the literature. For example, bispecific antibodies can be prepared using chemical linkage. Brennan et al., Science 229:81 (1985) describe a procedure

wherein intact antibodies are proteolytically cleaved to generate F(ab')<sub>2</sub> fragments. These fragments are reduced in the presence of the dithiol complexing agent sodium arsenite to stabilize vicinal dithiols and prevent intermolecular disulfide formation. The Fab' fragments generated are then converted to thionitrobenzoate (TNB) derivatives. One of the Fab'-TNB derivatives is then reconverted to the Fab'-thiol by reduction with mercaptoethylamine and is mixed with an equimolar amount of the other Fab'-TNB derivative to form the bispecific antibody. The bispecific antibodies produced can be used as agents for the selective immobilization of enzymes.

Additionally, Fab' fragments can be directly recovered from E. coli and chemically coupled to form bispecific antibodies. Shalaby et al., J. Exp. Med. 175:217-225 (1992) describe the production of a fully humanized bispecific antibody F(ab')<sub>2</sub> molecule. Each Fab' fragment was separately secreted from E. coli and subjected to directed chemical coupling in vitro to form the bispecific antibody. The bispecific antibody thus formed was able to bind to cells overexpressing the ErbB2 receptor and normal human T cells, as well as trigger the lytic activity of human cytotoxic lymphocytes against human breast tumor targets.

Various techniques for making and isolating bispecific antibody fragments directly from recombinant cell culture have also been described. For example, bispecific antibodies have been produced using leucine zippers. Kostelny et al., <u>J. Immunol.</u> 148(5):1547-1553 (1992). The leucine zipper peptides from the Fos and Jun proteins were linked to the Fab' portions of two different antibodies by gene fusion. The antibody homodimers were reduced at the hinge region to form monomers and then re-oxidized to form the antibody heterodimers. This method can also be utilized for the production of antibody homodimers. The "diabody" technology described by Hollinger et al., <u>Proc. Natl. Acad. Sci. USA</u> 90:6444-6448 (1993) has provided an alternative mechanism for making bispecific antibody fragments. The fragments comprise a heavy-chain variable domain (V<sub>H</sub>) connected to a light-chain variable domain (V<sub>L</sub>) by a linker which is too short to allow pairing between the two domains on the same chain. Accordingly, the V<sub>H</sub> and V<sub>L</sub> domains of one fragment are forced to pair with the complementary V<sub>L</sub> and V<sub>H</sub> domains of another fragment, thereby forming two antigen-binding sites. Another strategy for making bispecific antibody fragments by the use of single-chain Fv (sFv) dimers has also been reported. See, Gruber et al., <u>J. Immunol.</u> 152:5368 (1994).

Antibodies with more than two valencies are contemplated. For example, trispecific antibodies can be prepared. Tutt et al., <u>J. Immunol.</u> 147:60 (1991). Exemplary bispecific antibodies can bind to two different epitopes, at least one of which originates in the protein antigen of the invention. Alternatively, an anti-antigenic arm of an immunoglobulin molecule can be combined with an arm which binds to a triggering molecule on

a leukocyte such as a T-cell receptor molecule (e.g. CD2, CD3, CD28, or B7), or Fc receptors for IgG (Fc R), such as Fc RI (CD64), Fc RII (CD32) and Fc RIII (CD16) so as to focus cellular defense mechanisms to the cell expressing the particular antigen. Bispecific antibodies can also be used to direct cytotoxic agents to cells which express a particular antigen. These antibodies possess an antigen-binding arm and an arm which binds a cytotoxic agent or a radionuclide chelator, such as EOTUBE, DPTA, DOTA, or TETA. Another bispecific antibody of interest binds the protein antigen described herein and further binds tissue factor (TF).

# 5.13.6 Heteroconjugate Antibodies

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Heteroconjugate antibodies are also within the scope of the present invention.

Heteroconjugate antibodies are composed of two covalently joined antibodies. Such antibodies have, for example, been proposed to target immune system cells to unwanted cells (U.S. Patent No. 4,676,980), and for treatment of HIV infection (WO 91/00360; WO 92/200373; EP 03089). It is contemplated that the antibodies can be prepared in vitro using known methods in synthetic protein chemistry, including those involving crosslinking agents. For example, immunotoxins can be constructed using a disulfide exchange reaction or by forming a thioether bond. Examples of suitable reagents for this purpose include iminothiolate and methyl-4-mercaptobutyrimidate and those disclosed, for example, in U.S. Patent No. 4,676,980.

## 5.13.7 Effector Function Engineering

It can be desirable to modify the antibody of the invention with respect to effector function, so as to enhance, *e.g.*, the effectiveness of the antibody in treating cancer. For example, cysteine residue(s) can be introduced into the Fc region, thereby allowing interchain disulfide bond formation in this region. The homodimeric antibody thus generated can have improved internalization capability and/or increased complement-mediated cell killing and antibody-dependent cellular cytotoxicity (ADCC). See Caron et al., J. Exp Med., 176: 1191-1195 (1992) and Shopes, J. Immunol., 148: 2918-2922 (1992). Homodimeric antibodies with enhanced antitumor activity can also be prepared using heterobifunctional cross-linkers as described in Wolff et al. Cancer Research, 53: 2560-2565 (1993). Alternatively, an antibody can be engineered that has dual Fc regions and can thereby have enhanced complement lysis and ADCC capabilities. See Stevenson et al., Anti-Cancer Drug Design, 3: 219-230 (1989).

#### 5.13.8 Immunoconjugates

The invention also pertains to immunoconjugates comprising an antibody conjugated to a cytotoxic agent such as a chemotherapeutic agent, toxin (e.g., an enzymatically active toxin of

bacterial, fungal, plant, or animal origin, or fragments thereof), or a radioactive isotope (*i.e.*, a radioconjugate).

Chemotherapeutic agents useful in the generation of such immunoconjugates have been described above. Enzymatically active toxins and fragments thereof that can be used include diphtheria A chain, nonbinding active fragments of diphtheria toxin, exotoxin A chain (from Pseudomonas aeruginosa), ricin A chain, abrin A chain, modeccin A chain, alpha-sarcin, Aleurites fordii proteins, dianthin proteins, Phytolaca americana proteins (PAPI, PAPII, and PAP-S), momordica charantia inhibitor, curcin, crotin, sapaonaria officinalis inhibitor, gelonin, mitogellin, restrictocin, phenomycin, enomycin, and the tricothecenes. A variety of radionuclides are available for the production of radioconjugated antibodies. Examples include <sup>212</sup>Bi, <sup>131</sup>In, <sup>90</sup>Y, and <sup>186</sup>Re.

Conjugates of the antibody and cytotoxic agent are made using a variety of bifunctional protein-coupling agents such as N-succinimidyl-3-(2-pyridyldithiol) propionate (SPDP), iminothiolane (IT), bifunctional derivatives of imidoesters (such as dimethyl adipimidate HCL), active esters (such as disuccinimidyl suberate), aldehydes (such as glutareldehyde), bis-azido compounds (such as bis (p-azidobenzoyl) hexanediamine), bis-diazonium derivatives (such as bis-(p-diazoniumbenzoyl)-ethylenediamine), diisocyanates (such as tolyene 2,6-diisocyanate), and bis-active fluorine compounds (such as 1,5-difluoro-2,4-dinitrobenzene). For example, a ricin immunotoxin can be prepared as described in Vitetta et al., Science, 238: 1098 (1987). Carbon-14-labeled 1-isothiocyanatobenzyl-3-methyldiethylene triaminepentaacetic acid (MX-DTPA) is an exemplary chelating agent for conjugation of radionucleotide to the antibody. See WO94/11026.

In another embodiment, the antibody can be conjugated to a "receptor" (such streptavidin) for utilization in tumor pretargeting wherein the antibody-receptor conjugate is administered to the patient, followed by removal of unbound conjugate from the circulation using a clearing agent and then administration of a "ligand" (e.g., avidin) that is in turn conjugated to a cytotoxic agent.

# 4.14 COMPUTER READABLE SEQUENCES

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In one application of this embodiment, a nucleotide sequence of the present invention can be recorded on computer readable media. As used herein, "computer readable media" refers to any medium which can be read and accessed directly by a computer. Such media include, but are not limited to: magnetic storage media, such as floppy discs, hard disc storage medium, and magnetic tape; optical storage media such as CD-ROM; electrical storage media such as RAM and ROM; and hybrids of these categories such as magnetic/optical storage media. A skilled

artisan can readily appreciate how any of the presently known computer readable mediums can be used to create a manufacture comprising computer readable medium having recorded thereon a nucleotide sequence of the present invention. As used herein, "recorded" refers to a process for storing information on computer readable medium. A skilled artisan can readily adopt any of the presently known methods for recording information on computer readable medium to generate manufactures comprising the nucleotide sequence information of the present invention.

A variety of data storage structures are available to a skilled artisan for creating a computer readable medium having recorded thereon a nucleotide sequence of the present invention. The choice of the data storage structure will generally be based on the means chosen to access the stored information. In addition, a variety of data processor programs and formats can be used to store the nucleotide sequence information of the present invention on computer readable medium. The sequence information can be represented in a word processing text file, formatted in commercially-available software such as WordPerfect and Microsoft Word, or represented in the form of an ASCII file, stored in a database application, such as DB2, Sybase, Oracle, or the like. A skilled artisan can readily adapt any number of data processor structuring formats (e.g. text file or database) in order to obtain computer readable medium having recorded thereon the nucleotide sequence information of the present invention.

By providing any of the nucleotide sequences SEQ ID NO:1-1350 or a representative fragment thereof; or a nucleotide sequence at least 95% identical to any of the nucleotide sequences of SEQ ID NO:1-1350 in computer readable form, a skilled artisan can routinely access the sequence information for a variety of purposes. Computer software is publicly available which allows a skilled artisan to access sequence information provided in a computer readable medium. The examples which follow demonstrate how software which implements the BLAST (Altschul et al., J. Mol. Biol. 215:403-410 (1990)) and BLAZE (Brutlag et al., Comp. Chem. 17:203-207 (1993)) search algorithms on a Sybase system is used to identify open reading frames (ORFs) within a nucleic acid sequence. Such ORFs may be protein encoding fragments and may be useful in producing commercially important proteins such as enzymes used in fermentation reactions and in the production of commercially useful metabolites.

As used herein, "a computer-based system" refers to the hardware means, software means, and data storage means used to analyze the nucleotide sequence information of the present invention. The minimum hardware means of the computer-based systems of the present invention comprises a central processing unit (CPU), input means, output means, and data storage means. A skilled artisan can readily appreciate that any one of the currently available computer-based systems are suitable for use in the present invention. As stated above, the computer-based systems of the present invention comprise a data storage means having stored

therein a nucleotide sequence of the present invention and the necessary hardware means and software means for supporting and implementing a search means. As used herein, "data storage means" refers to memory which can store nucleotide sequence information of the present invention, or a memory access means which can access manufactures having recorded thereon the nucleotide sequence information of the present invention.

As used herein, "search means" refers to one or more programs which are implemented on the computer-based system to compare a target sequence or target structural motif with the sequence information stored within the data storage means. Search means are used to identify fragments or regions of a known sequence which match a particular target sequence or target motif. A variety of known algorithms are disclosed publicly and a variety of commercially available software for conducting search means are and can be used in the computer-based systems of the present invention. Examples of such software includes, but is not limited to, Smith-Waterman, MacPattern (EMBL), BLASTN and BLASTA (NPOLYPEPTIDEIA). A skilled artisan can readily recognize that any one of the available algorithms or implementing software packages for conducting homology searches can be adapted for use in the present computer-based systems. As used herein, a "target sequence" can be any nucleic acid or amino acid sequence of six or more nucleotides or two or more amino acids. A skilled artisan can readily recognize that the longer a target sequence is, the less likely a target sequence will be present as a random occurrence in the database. The most preferred sequence length of a target sequence is from about 10 to 300 amino acids, more preferably from about 30 to 100 nucleotide residues. However, it is well recognized that searches for commercially important fragments, such as sequence fragments involved in gene expression and protein processing, may be of shorter length.

As used herein, "a target structural motif," or "target motif," refers to any rationally selected sequence or combination of sequences in which the sequence(s) are chosen based on a three-dimensional configuration which is formed upon the folding of the target motif. There are a variety of target motifs known in the art. Protein target motifs include, but are not limited to, enzyme active sites and signal sequences. Nucleic acid target motifs include, but are not limited to, promoter sequences, hairpin structures and inducible expression elements (protein binding sequences).

## 4.15 TRIPLE HELIX FORMATION

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In addition, the fragments of the present invention, as broadly described, can be used to control gene expression through triple helix formation or antisense DNA or RNA, both of which methods are based on the binding of a polynucleotide sequence to DNA or RNA.

Polynucleotides suitable for use in these methods are preferably 20 to 40 bases in length and are designed to be complementary to a region of the gene involved in transcription (triple helix - see Lee et al., Nucl. Acids Res. 6:3073 (1979); Cooney et al., Science 15241:456 (1988); and Dervan et al., Science 251:1360 (1991)) or to the mRNA itself (antisense - Olmno, J. Neurochem. 56:560 (1991); Oligodeoxynucleotides as Antisense Inhibitors of Gene Expression, CRC Press, Boca Raton, FL (1988)). Triple helix-formation optimally results in a shut-off of RNA transcription from DNA, while antisense RNA hybridization blocks translation of an mRNA molecule into polypeptide. Both techniques have been demonstrated to be effective in model systems. Information contained in the sequences of the present invention is necessary for the design of an antisense or triple helix oligonucleotide.

## 4.16 DIAGNOSTIC ASSAYS AND KITS

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The present invention further provides methods to identify the presence or expression of one of the ORFs of the present invention, or homolog thereof, in a test sample, using a nucleic acid probe or antibodies of the present invention, optionally conjugated or otherwise associated with a suitable label.

In general, methods for detecting a polynucleotide of the invention can comprise contacting a sample with a compound that binds to and forms a complex with the polynucleotide for a period sufficient to form the complex, and detecting the complex, so that if a complex is detected, a polynucleotide of the invention is detected in the sample. Such methods can also comprise contacting a sample under stringent hybridization conditions with nucleic acid primers that anneal to a polynucleotide of the invention under such conditions, and amplifying annealed polynucleotides, so that if a polynucleotide is amplified, a polynucleotide of the invention is detected in the sample.

In general, methods for detecting a polypeptide of the invention can comprise contacting a sample with a compound that binds to and forms a complex with the polypeptide for a period sufficient to form the complex, and detecting the complex, so that if a complex is detected, a polypeptide of the invention is detected in the sample.

In detail, such methods comprise incubating a test sample with one or more of the antibodies or one or more of the nucleic acid probes of the present invention and assaying for binding of the nucleic acid probes or antibodies to components within the test sample.

Conditions for incubating a nucleic acid probe or antibody with a test sample vary. Incubation conditions depend on the format employed in the assay, the detection methods employed, and the type and nature of the nucleic acid probe or antibody used in the assay. One skilled in the art will recognize that any one of the commonly available hybridization,

amplification or immunological assay formats can readily be adapted to employ the nucleic acid probes or antibodies of the present invention. Examples of such assays can be found in Chard, T., An Introduction to Radioimmunoassay and Related Techniques, Elsevier Science Publishers, Amsterdam, The Netherlands (1986); Bullock, G.R. et al., Techniques in Immunocytochemistry, Academic Press, Orlando, FL Vol. 1 (1982), Vol. 2 (1983), Vol. 3 (1985); Tijssen, P., Practice and Theory of immunoassays: Laboratory Techniques in Biochemistry and Molecular Biology, Elsevier Science Publishers, Amsterdam, The Netherlands (1985). The test samples of the present invention include cells, protein or membrane extracts of cells, or biological fluids such as sputum, blood, serum, plasma, or urine. The test sample used in the above-described method will vary based on the assay format, nature of the detection method and the tissues, cells or extracts used as the sample to be assayed. Methods for preparing protein extracts or membrane extracts of cells are well known in the art and can be readily be adapted in order to obtain a sample which is compatible with the system utilized.

In another embodiment of the present invention, kits are provided which contain the necessary reagents to carry out the assays of the present invention. Specifically, the invention provides a compartment kit to receive, in close confinement, one or more containers which comprises: (a) a first container comprising one of the probes or antibodies of the present invention; and (b) one or more other containers comprising one or more of the following: wash reagents, reagents capable of detecting presence of a bound probe or antibody.

In detail, a compartment kit includes any kit in which reagents are contained in separate containers. Such containers include small glass containers, plastic containers or strips of plastic or paper. Such containers allows one to efficiently transfer reagents from one compartment to another compartment such that the samples and reagents are not cross-contaminated, and the agents or solutions of each container can be added in a quantitative fashion from one compartment to another. Such containers will include a container which will accept the test sample, a container which contains the antibodies used in the assay, containers which contain wash reagents (such as phosphate buffered saline, Tris-buffers, etc.), and containers which contain the reagents used to detect the bound antibody or probe. Types of detection reagents include labeled nucleic acid probes, labeled secondary antibodies, or in the alternative, if the primary antibody is labeled, the enzymatic, or antibody binding reagents which are capable of reacting with the labeled antibody. One skilled in the art will readily recognize that the disclosed probes and antibodies of the present invention can be readily incorporated into one of the established kit formats which are well known in the art.

## 4.17 MEDICAL IMAGING

The novel polypeptides and binding partners of the invention are useful in medical imaging of sites expressing the molecules of the invention (e.g., where the polypeptide of the invention is involved in the immune response, for imaging sites of inflammation or infection). See, e.g., Kunkel et al., U.S. Pat. NO. 5,413,778. Such methods involve chemical attachment of a labeling or imaging agent, administration of the labeled polypeptide to a subject in a pharmaceutically acceptable carrier, and imaging the labeled polypeptide in vivo at the target site.

#### 4.18 SCREENING ASSAYS

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Using the isolated proteins and polynucleotides of the invention, the present invention further provides methods of obtaining and identifying agents which bind to a polypeptide encoded by an ORF corresponding to any of the nucleotide sequences set forth in SEQ ID NO:1-1350, or bind to a specific domain of the polypeptide encoded by the nucleic acid. In detail, said method comprises the steps of:

- (a) contacting an agent with an isolated protein encoded by an ORF of the present invention, or nucleic acid of the invention; and
  - (b) determining whether the agent binds to said protein or said nucleic acid.

In general, therefore, such methods for identifying compounds that bind to a polynucleotide of the invention can comprise contacting a compound with a polynucleotide of the invention for a time sufficient to form a polynucleotide/compound complex, and detecting the complex, so that if a polynucleotide/compound complex is detected, a compound that binds to a polynucleotide of the invention is identified.

Likewise, in general, therefore, such methods for identifying compounds that bind to a polypeptide of the invention can comprise contacting a compound with a polypeptide of the invention for a time sufficient to form a polypeptide/compound complex, and detecting the complex, so that if a polypeptide/compound complex is detected, a compound that binds to a polynucleotide of the invention is identified.

Methods for identifying compounds that bind to a polypeptide of the invention can also comprise contacting a compound with a polypeptide of the invention in a cell for a time sufficient to form a polypeptide/compound complex, wherein the complex drives expression of a receptor gene sequence in the cell, and detecting the complex by detecting reporter gene sequence expression, so that if a polypeptide/compound complex is detected, a compound that binds a polypeptide of the invention is identified.

Compounds identified via such methods can include compounds which modulate the activity of a polypeptide of the invention (that is, increase or decrease its activity, relative to

activity observed in the absence of the compound). Alternatively, compounds identified via such methods can include compounds which modulate the expression of a polynucleotide of the invention (that is, increase or decrease expression relative to expression levels observed in the absence of the compound). Compounds, such as compounds identified via the methods of the invention, can be tested using standard assays well known to those of skill in the art for their ability to modulate activity/expression.

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The agents screened in the above assay can be, but are not limited to, peptides, carbohydrates, vitamin derivatives, or other pharmaceutical agents. The agents can be selected and screened at random or rationally selected or designed using protein modeling techniques.

For random screening, agents such as peptides, carbohydrates, pharmaceutical agents and the like are selected at random and are assayed for their ability to bind to the protein encoded by the ORF of the present invention. Alternatively, agents may be rationally selected or designed. As used herein, an agent is said to be "rationally selected or designed" when the agent is chosen based on the configuration of the particular protein. For example, one skilled in the art can readily adapt currently available procedures to generate peptides, pharmaceutical agents and the like, capable of binding to a specific peptide sequence, in order to generate rationally designed antipeptide peptides, for example see Hurby et al., Application of Synthetic Peptides: Antisense Peptides," In Synthetic Peptides, A User's Guide, W.H. Freeman, NY (1992), pp. 289-307, and Kaspczak et al., Biochemistry 28:9230-8 (1989), or pharmaceutical agents, or the like.

In addition to the foregoing, one class of agents of the present invention, as broadly described, can be used to control gene expression through binding to one of the ORFs or EMFs of the present invention. As described above, such agents can be randomly screened or rationally designed/selected. Targeting the ORF or EMF allows a skilled artisan to design sequence specific or element specific agents, modulating the expression of either a single ORF or multiple ORFs which rely on the same EMF for expression control. One class of DNA binding agents are agents which contain base residues which hybridize or form a triple helix formation by binding to DNA or RNA. Such agents can be based on the classic phosphodiester, ribonucleic acid backbone, or can be a variety of sulfhydryl or polymeric derivatives which have base attachment capacity.

Agents suitable for use in these methods preferably contain 20 to 40 bases and are designed to be complementary to a region of the gene involved in transcription (triple helix - see Lee et al., Nucl. Acids Res. 6:3073 (1979); Cooney et al., Science 241:456 (1988); and Dervan et al., Science 251:1360 (1991)) or to the mRNA itself (antisense - Okano, J. Neurochem. 56:560 (1991); Oligodeoxynucleotides as Antisense Inhibitors of Gene Expression, CRC Press, Boca Raton, FL (1988)). Triple helix-formation optimally results in a shut-off of RNA transcription

from DNA, while antisense RNA hybridization blocks translation of an mRNA molecule into polypeptide. Both techniques have been demonstrated to be effective in model systems. Information contained in the sequences of the present invention is necessary for the design of an antisense or triple helix oligonucleotide and other DNA binding agents.

Agents which bind to a protein encoded by one of the ORFs of the present invention can be used as a diagnostic agent. Agents which bind to a protein encoded by one of the ORFs of the present invention can be formulated using known techniques to generate a pharmaceutical composition.

## 4.19 USE OF NUCLEIC ACIDS AS PROBES

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Another aspect of the subject invention is to provide for polypeptide-specific nucleic acid hybridization probes capable of hybridizing with naturally occurring nucleotide sequences. The hybridization probes of the subject invention may be derived from any of the nucleotide sequences SEQ ID NO:1-1350. Because the corresponding gene is only expressed in a limited number of tissues, a hybridization probe derived from of any of the nucleotide sequences SEQ ID NO:1-1350 can be used as an indicator of the presence of RNA of cell type of such a tissue in a sample.

Any suitable hybridization technique can be employed, such as, for example, in situ hybridization. PCR as described in US Patents Nos. 4,683,195 and 4,965,188 provides additional uses for oligonucleotides based upon the nucleotide sequences. Such probes used in PCR may be of recombinant origin, may be chemically synthesized, or a mixture of both. The probe will comprise a discrete nucleotide sequence for the detection of identical sequences or a degenerate pool of possible sequences for identification of closely related genomic sequences.

Other means for producing specific hybridization probes for nucleic acids include the cloning of nucleic acid sequences into vectors for the production of mRNA probes. Such vectors are known in the art and are commercially available and may be used to synthesize RNA probes in vitro by means of the addition of the appropriate RNA polymerase as T7 or SP6 RNA polymerase and the appropriate radioactively labeled nucleotides. The nucleotide sequences may be used to construct hybridization probes for mapping their respective genomic sequences. The nucleotide sequence provided herein may be mapped to a chromosome or specific regions of a chromosome using well known genetic and/or chromosomal mapping techniques. These techniques include in situ hybridization, linkage analysis against known chromosomal markers, hybridization screening with libraries or flow-sorted chromosomal preparations specific to known chromosomes, and the like. The technique of fluorescent in situ hybridization of

chromosome spreads has been described, among other places, in Verma et al (1988) Human Chromosomes: A Manual of Basic Techniques, Pergamon Press, New York NY.

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Fluorescent *in situ* hybridization of chromosomal preparations and other physical chromosome mapping techniques may be correlated with additional genetic map data. Examples of genetic map data can be found in the 1994 Genome Issue of Science (265:1981f). Correlation between the location of a nucleic acid on a physical chromosomal map and a specific disease (or predisposition to a specific disease) may help delimit the region of DNA associated with that genetic disease. The nucleotide sequences of the subject invention may be used to detect differences in gene sequences between normal, carrier or affected individuals.

## 4.20 PREPARATION OF SUPPORT BOUND OLIGONUCLEOTIDES

Oligonucleotides, *i.e.*, small nucleic acid segments, may be readily prepared by, for example, directly synthesizing the oligonucleotide by chemical means, as is commonly practiced using an automated oligonucleotide synthesizer.

Support bound oligonucleotides may be prepared by any of the methods known to those of skill in the art using any suitable support such as glass, polystyrene or Teflon. One strategy is to precisely spot oligonucleotides synthesized by standard synthesizers. Immobilization can be achieved using passive adsorption (Inouye & Hondo, (1990) J. Clin. Microbiol. 28(6) 1469-72); using UV light (Nagata *et al.*, 1985; Dahlen *et al.*, 1987; Morrissey & Collins, (1989) Mol. Cell Probes 3(2) 189-207) or by covalent binding of base modified DNA (Keller *et al.*, 1988; 1989); all references being specifically incorporated herein.

Another strategy that may be employed is the use of the strong biotin-streptavidin interaction as a linker. For example, Broude *et al.* (1994) Proc. Natl. Acad. Sci. USA 91(8) 3072-6, describe the use of biotinylated probes, although these are duplex probes, that are immobilized on streptavidin-coated magnetic beads. Streptavidin-coated beads may be purchased from Dynal, Oslo. Of course, this same linking chemistry is applicable to coating any surface with streptavidin. Biotinylated probes may be purchased from various sources, such as, *e.g.*, Operon Technologies (Alameda, CA).

Nunc Laboratories (Naperville, IL) is also selling suitable material that could be used. Nunc Laboratories have developed a method by which DNA can be covalently bound to the microwell surface termed Covalink NH. CovaLink NH is a polystyrene surface grafted with secondary amino groups (>NH) that serve as bridge-heads for further covalent coupling. CovaLink Modules may be purchased from Nunc Laboratories. DNA molecules may be bound to CovaLink exclusively at the 5'-end by a phosphoramidate bond, allowing immobilization of more than 1 pmol of DNA (Rasmussen *et al.*, (1991) Anal. Biochem. 198(1) 138-42).

The use of CovaLink NH strips for covalent binding of DNA molecules at the 5'-end has been described (Rasmussen et al., (1991). In this technology, a phosphoramidate bond is employed (Chu et al., (1983) Nucleic Acids Res. 11(8) 6513-29). This is beneficial as immobilization using only a single covalent bond is preferred. The phosphoramidate bond joins the DNA to the CovaLink NH secondary amino groups that are positioned at the end of spacer arms covalently grafted onto the polystyrene surface through a 2 nm long spacer arm. To link an oligonucleotide to CovaLink NH via an phosphoramidate bond, the oligonucleotide terminus must have a 5'-end phosphate group. It is, perhaps, even possible for biotin to be covalently bound to CovaLink and then streptavidin used to bind the probes.

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More specifically, the linkage method includes dissolving DNA in water (7.5 ng/ul) and denaturing for 10 min. at 95°C and cooling on ice for 10 min. Ice-cold 0.1 M 1-methylimidazole, pH 7.0 (1-MeIm<sub>7</sub>), is then added to a final concentration of 10 mM 1-MeIm<sub>7</sub>. A ss DNA solution is then dispensed into CovaLink NH strips (75 ul/well) standing on ice.

Carbodiimide 0.2 M 1-ethyl-3-(3-dimethylaminopropyl)-carbodiimide (EDC), dissolved in 10 mM 1-MeIm<sub>7</sub>, is made fresh and 25 ul added per well. The strips are incubated for 5 hours at 50°C. After incubation the strips are washed using, *e.g.*, Nunc-Immuno Wash; first the wells are washed 3 times, then they are soaked with washing solution for 5 min., and finally they are washed 3 times (where in the washing solution is 0.4 N NaOH, 0.25% SDS heated to 50°C).

It is contemplated that a further suitable method for use with the present invention is that described in PCT Patent Application WO 90/03382 (Southern & Maskos), incorporated herein by reference. This method of preparing an oligonucleotide bound to a support involves attaching a nucleoside 3'-reagent through the phosphate group by a covalent phosphodiester link to aliphatic hydroxyl groups carried by the support. The oligonucleotide is then synthesized on the supported nucleoside and protecting groups removed from the synthetic oligonucleotide chain under standard conditions that do not cleave the oligonucleotide from the support. Suitable reagents include nucleoside phosphoramidite and nucleoside hydrogen phosphorate.

An on-chip strategy for the preparation of DNA probe for the preparation of DNA probe arrays may be employed. For example, addressable laser-activated photodeprotection may be employed in the chemical synthesis of oligonucleotides directly on a glass surface, as described by Fodor *et al.* (1991) Science 251(4995) 767-73, incorporated herein by reference. Probes may also be immobilized on nylon supports as described by Van Ness *et al.* (1991) Nucleic Acids Res. 19(12) 3345-50; or linked to Teflon using the method of Duncan & Cavalier (1988) Anal. Biochem. 169(1) 104-8; all references being specifically incorporated herein.

To link an oligonucleotide to a nylon support, as described by Van Ness *et al.* (1991), requires activation of the nylon surface via alkylation and selective activation of the 5'-amine of oligonucleotides with cyanuric chloride.

One particular way to prepare support bound oligonucleotides is to utilize the light-generated synthesis described by Pease *et al.*, (1994) PNAS USA 91(11) 5022-6, incorporated herein by reference). These authors used current photolithographic techniques to generate arrays of immobilized oligonucleotide probes (DNA chips). These methods, in which light is used to direct the synthesis of oligonucleotide probes in high-density, miniaturized arrays, utilize photolabile 5'-protected *N*-acyl-deoxynucleoside phosphoramidites, surface linker chemistry and versatile combinatorial synthesis strategies. A matrix of 256 spatially defined oligonucleotide probes may be generated in this manner.

## 4.21 PREPARATION OF NUCLEIC ACID FRAGMENTS

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The nucleic acids may be obtained from any appropriate source, such as cDNAs, genomic DNA, chromosomal DNA, microdissected chromosome bands, cosmid or YAC inserts, and RNA, including mRNA without any amplification steps. For example, Sambrook *et al.* (1989) describes three protocols for the isolation of high molecular weight DNA from mammalian cells (p. 9.14-9.23).

DNA fragments may be prepared as clones in M13, plasmid or lambda vectors and/or prepared directly from genomic DNA or cDNA by PCR or other amplification methods. Samples may be prepared or dispensed in multiwell plates. About 100-1000 ng of DNA samples may be prepared in 2-500 ml of final volume.

The nucleic acids would then be fragmented by any of the methods known to those of skill in the art including, for example, using restriction enzymes as described at 9.24-9.28 of Sambrook *et al.* (1989), shearing by ultrasound and NaOH treatment.

Low pressure shearing is also appropriate, as described by Schriefer *et al.* (1990) Nucleic Acids Res. 18(24) 7455-6, incorporated herein by reference). In this method, DNA samples are passed through a small French pressure cell at a variety of low to intermediate pressures. A lever device allows controlled application of low to intermediate pressures to the cell. The results of these studies indicate that low-pressure shearing is a useful alternative to sonic and enzymatic DNA fragmentation methods.

One particularly suitable way for fragmenting DNA is contemplated to be that using the two base recognition endonuclease, *Cvi*JI, described by Fitzgerald *et al.* (1992) Nucleic Acids Res. 20(14) 3753-62. These authors described an approach for the rapid fragmentation and fractionation

of DNA into particular sizes that they contemplated to be suitable for shotgun cloning and sequencing.

The restriction endonuclease *Cvi*JI normally cleaves the recognition sequence PuGCPy between the G and C to leave blunt ends. Atypical reaction conditions, which alter the specificity of this enzyme (*Cvi*JI\*\*), yield a quasi-random distribution of DNA fragments form the small molecule pUC19 (2688 base pairs). Fitzgerald *et al.* (1992) quantitatively evaluated the randomness of this fragmentation strategy, using a *Cvi*JI\*\* digest of pUC19 that was size fractionated by a rapid gel filtration method and directly ligated, without end repair, to a lac Z minus M13 cloning vector. Sequence analysis of 76 clones showed that *Cvi*JI\*\* restricts pyGCPy and PuGCPu, in addition to PuGCPy sites, and that new sequence data is accumulated at a rate consistent with random fragmentation.

As reported in the literature, advantages of this approach compared to sonication and agarose gel fractionation include: smaller amounts of DNA are required (0.2-0.5 ug instead of 2-5 ug); and fewer steps are involved (no preligation, end repair, chemical extraction, or agarose gel electrophoresis and elution are needed

Irrespective of the manner in which the nucleic acid fragments are obtained or prepared, it is important to denature the DNA to give single stranded pieces available for hybridization. This is achieved by incubating the DNA solution for 2-5 minutes at 80-90°C. The solution is then cooled quickly to 2°C to prevent renaturation of the DNA fragments before they are contacted with the chip. Phosphate groups must also be removed from genomic DNA by methods known in the art.

#### 4.22 PREPARATION OF DNA ARRAYS

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Arrays may be prepared by spotting DNA samples on a support such as a nylon membrane. Spotting may be performed by using arrays of metal pins (the positions of which correspond to an array of wells in a microtiter plate) to repeated by transfer of about 20 nl of a DNA solution to a nylon membrane. By offset printing, a density of dots higher than the density of the wells is achieved. One to 25 dots may be accommodated in 1 mm², depending on the type of label used. By avoiding spotting in some preselected number of rows and columns, separate subsets (subarrays) may be formed. Samples in one subarray may be the same genomic segment of DNA (or the same gene) from different individuals, or may be different, overlapped genomic clones. Each of the subarrays may represent replica spotting of the same samples. In one example, a selected gene segment may be amplified from 64 patients. For each patient, the amplified gene segment may be in one 96-well plate (all 96 wells containing the same sample). A plate for each of the 64 patients is prepared. By using a 96-pin device, all samples may be spotted on one 8 x 12 cm membrane.

Subarrays may contain 64 samples, one from each patient. Where the 96 subarrays are identical, the dot span may be 1 mm<sup>2</sup> and there may be a 1 mm space between subarrays.

Another approach is to use membranes or plates (available from NUNC, Naperville, Illinois) which may be partitioned by physical spacers *e.g.* a plastic grid molded over the membrane, the grid being similar to the sort of membrane applied to the bottom of multiwell plates, or hydrophobic strips. A fixed physical spacer is not preferred for imaging by exposure to flat phosphor-storage screens or x-ray films.

The present invention is illustrated in the following examples. Upon consideration of the present disclosure, one of skill in the art will appreciate that many other embodiments and variations may be made in the scope of the present invention. Accordingly, it is intended that the broader aspects of the present invention not be limited to the disclosure of the following examples. The present invention is not to be limited in scope by the exemplified embodiments which are intended as illustrations of single aspects of the invention, and compositions and methods which are functionally equivalent are within the scope of the invention. Indeed, numerous modifications and variations in the practice of the invention are expected to occur to those skilled in the art upon consideration of the present preferred embodiments. Consequently, the only limitations which should be placed upon the scope of the invention are those which appear in the appended claims.

All references cited within the body of the instant specification are hereby incorporated by reference in their entirety.

## 5.0 EXAMPLES

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#### 5.1 EXAMPLE 1

## Novel Nucleic Acid Sequences Obtained From Various Libraries

A plurality of novel nucleic acids were obtained from cDNA libraries prepared from various human tissues and in some cases isolated from a genomic library derived from human chromosome using standard PCR, SBH sequence signature analysis and Sanger sequencing techniques. The inserts of the library were amplified with PCR using primers specific for the vector sequences which flank the inserts. Clones from cDNA libraries were spotted on nylon membrane filters and screened with oligonucleotide probes (e.g., 7-mers) to obtain signature sequences. The clones were clustered into groups of similar or identical sequences. Representative clones were selected for sequencing.

In some cases, the 5' sequence of the amplified inserts was then deduced using a typical Sanger sequencing protocol. PCR products were purified and subjected to fluorescent dye terminator cycle sequencing. Single pass gel sequencing was done using a 377 Applied Biosystems

(ABI) sequencer to obtain the novel nucleic acid sequences. In some cases RACE (Random Amplification of cDNA Ends) was performed to further extend the sequence in the 5' direction.

#### 5.2 EXAMPLE 2

# 5 Novel Contigs

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The novel contigs of the invention were assembled from sequences that were obtained from a cDNA library by methods described in Example 1 above, and in some cases sequences obtained from one or more public databases. The sequences for the resulting nucleic acid contigs are designated as SEQ ID NO: 1-1350 and are provided in the attached Sequence Listing. The contigs were assembled using an EST sequence as a seed. Then a recursive algorithm was used to extend the seed EST into an extended assemblage, by pulling additional sequences from different databases (i.e., Hyseq's database containing EST sequences, dbEST version 114, gb pri 114, and UniGene version 101) that belong to this assemblage. The algorithm terminated when there was no additional sequences from the above databases that would extend the assemblage. Inclusion of component sequences into the assemblage was based on a BLASTN hit to the extending assemblage with BLAST score greater than 300 and percent identity greater than 95%.

Table 3 sets forth the novel predicted polypeptides (including proteins) encoded by the novel polynucleotides (SEQ ID NO:189-282) of the present invention, and their corresponding nucleotide locations to each of SEQ ID NO: 189-282. Table 3 also indicates the method by which the polypeptide was predicted. Method A refers to a polypeptide obtained by using a software program called FASTY (available from <a href="http://fasta.bioch.virginia.edu">http://fasta.bioch.virginia.edu</a>) which selects a polypeptide based on a comparison of the translated novel polynucleotide to known polynucleotides (W.R. Pearson, Methods in Enzymology, 183:63-98 (1990), herein incorporated by reference). Method B refers to a polypeptide obtained by using a software program called GenScan for human/vertebrate sequences (available from Stanford University, Office of Technology Licensing) that predicts the polypeptide based on a probabilistic model of gene structure/compositional properties (C. Burge and S. Karlin, J. Mol. Biol., 268:78-94 (1997), incorporated herein by reference). Method C refers to a polypeptide obtained by using a Hyseq proprietary software program that translates the novel polynucleotide and its complementary strand into six possible amino acid sequences (forward and reverse frames) and chooses the polypeptide with the longest open reading frame.

The nearest neighbor results for SEQ ID NO: 1-1350 were obtained by a BLASTP version 2.0al 19MP-WashU search against Genpept release 120 and Geneseq database October 12, 2000, update 21 (Derwent), using BLAST algorithm. The nearest neighbor result showed the

closest homologue for SEQ ID NO:1-1350. The nearest neighbor results for SEQ ID NO: 1-1350 are shown in Table 2 below.

Tables 1, 2 and 3 follow. Table 1 shows the various tissue sources of SEQ ID NO: 1-1350. Table 2 shows the nearest neighbor result for the assembled contig. The nearest neighbor result shows the closest homolog with an identifiable function for each assemblage. Table 3 contains the start and stop nucleotides for the translated amino acid sequence for which each assemblage encodes. Table 3 also provides a correlation between the amino acid sequences set forth in the Sequence Listing, the nucleotide sequences set forth in the Sequence Listing and the SEQ ID NO. in USSN 09/496,914.

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# TABLE 1

Tissue Origin	RNA Source	Hyseq Library Name	SEQ ID NOS:
adult brain	GIBCO	AB3001	111 151 188 215 662-665 877 910 927
	j		976 1233 1319
adult brain	GIBCO	ABD003	41 49 74 101 111 120 132 141-142 151
			217 225 238 271 317 404 446 469 503
		!	513-514 535 550 564 573 666-669 798
1		1	898 910 927 976 1067 1083 1085 1178
			1254
adult brain	Clontech	ABR001	39 216 238 327 356 535 927 1056 1121
1 1(1 '	<u> </u>	4 7 7 0 0 ¢	1178-1180 1199 1251
adult brain	Clontech	ABR006	74 611 949 1034 1136
adult brain	Clontech	ABR008	14 32 41 61 81 86 89 120 132 138 145
		•	147 188 197 208 225 227-239 250 300-
			303 312 316 328-331 340 357-362 374 380 384-391 408 414 446 448 464-467
			483 488 495-496 505 512 521 535 550
		ļ	566 571 577 585 590 594 598 634 641
			658 666 683 725 742 764 767 786 801
			805 810 823 826 829 831 836 841 887-
			923 927 934 943 950-951 963 976 995
			1000-1001 1006 1026 1034 1048 1057-
		Ì	1067 1086 1088 1090 1118 1120 1122-
		ļ.	1128 1142 1162 1181-1192 1199 1204
			1218-1219 1225 1232 1253 1267 1271-
			1306 1342 1347 1349-1350
adult brain	Clontech	ABR011	49 238 1219
adult brain	BioChain	ABR012	74 238
adult brain	Invitrogen	ABR013	868 1268
adult brain	Invitrogen	ABT004	49 117 138 191 217 252 291 305 535
			566 596 663 670 746 798 816-819 876
			892 898 922 943 963 1034-1036 1121
cultured	Strategene	ADP001	41 74 101 138 211 238 304 537 582
preadipocytes		1777000	740 798 883 943 976 1067
adrenal gland	Clontech	ADR002	49 74 101 111 120 127 151 215 238
			240-247 316 330 363-364 404 414 534-
			535 833 924-940 950 963 976 1001
			1325
adult heart	GIBCO	AHR001	38 49 71-72 74-77 79 92 99 101 111
uddit Hourt	GIBCO	THIROUT	118 129 132 138 151 158-163 182 195-
			203 215 217 238 264 269 353 384 398
			408 434-439 446 504 512-513 519 537
			562-573 577 611-614 616-619 658 661
	į		671-672 722 734 757-773 815 828-835
			874 891 898 919 926-927 976 988
			1021 1037 1041 1062 1067 1071 1080
			1083 1093 1122 1131 1185 1201 1254
			1308 1331 1335
adult kidney	GIBCO	AKD001	41 49 51 71-74 78-85 94 100-101 103-
	{		107 111 119-120 138 151 157 215 217-
			218 238 250 264 294 304 384 404 440
	}		446 454 477 504-505 509 514 518-519
			535 537 564 574-583 620-627 639 653
			673-675 705 753 789 831 844 851 859
			877 909 918 927 956 963 976 1067
adult kidney	Invitoocc	AKTOO	1074 1083 1095 1178 1302 1331 1335
addit vidileh	Invitrogen	AKT002	11-12 41 49 111-112 215-217 294 316 446 487 564 575 844 868 910 927 976
			1116
adult lung	GIBCO	ALG001	8 101 111 151 187 402 446 490 514
addit fullg	- GIDCO	LVEGOOT	0 101 111 131 107 402 440 430 314

Tissue Origin	RNA Source	Hyseq Library N	ame   SEQ ID NOS:
			518 537 545 549 580 582 592 594 634
		1	640 651-652 676-678 725 851 873 918
			952 976 1042 1067 1076 1083 1152
lymph node	Clontech	ALN001	8 111 121 151 180-182 188 215 537
			545 549 651 679-682 789 804-810 868
			873 927 952 976 1042 1059 1335
young liver	GIBCO	ALV001	8 64 79 111 186 215-216 238 446 514
			519 537 564 653 683-684 698 753 798
			813 833 840 858 927 976 1038-1039
			1051 1085 1224 1245 1256
adult liver	Invitrogen	ALV002	40 71 292-293 305 384 468-469 496
	Ì		505 657 675 714 753 832 844 941-942
			976 1040 1076 1256 1293
adult liver	Clontech	ALV003	976
adult ovary	Invitrogen	AOV001	8 32 36 38 41 49 51 71 74 79-80 101
	}		104 111 120 122-125 138 140 143-149
			151 188-190 207-212 215-217 238 264
			316 384 409 440 445-446 496 504 512
			514 518-519 535 537 549-550 564 566
			571 580 582 600 618 638 657 667 681
			685-697 699 705 722 735-744 761 771
			815 833 842-865 868 875-876 918 926-
			927 950 952 963 976 1023 1042 1048
			1051 1059 1072 1076 1083 1117 1120
			1124 1131 1144 1174 1224 1268 1331
			1335
adult placenta	Clontech	APL001	102 217 238 537 641 700
placenta	Invitrogen	APL002	663 851 1048
adult spleen	GIBCO	ASP001	8 45 74 111 132 140 151 185 217 238
			294 414 446 477 504 514 534 545 549
			592 722 873 883 952 976 1041-1042
A. 1.	- CTRCC	A75001	1083 1093-1094 1152 1224
testis	GIBCO	ATS001	72 107 111 113 126 140 151 183 215
			238 446 497 537 642 701-706 811 877
111111		DI 12001	927 962 976 1083 1117 1131
adult bladder	Invitrogen	BLD001	41 151 191 402-405 409 414 496 545
1	01	DMD001	592 607 706 873 952 1178 1329-1335
bone marrow	Clontech	BMD001	8 58-62 65-68 74 79 108 111 116 137
			147 151 164-174 213-215 238 305-307
			374 404 446 460 466 516 519 534 538- 541 544-546 549-554 566 584 586 592
	•		
			596 607 610 628-629 643-645 652 707- 708 774-789 844 866-871 873 919 927
	1		952 963 976 998 1034 1042 1064 1083
			1085 1120 1132 1152 1225 1229 1268
	İ		1307 1310
bone marrow	Clontech	BMD002	6 8 37-38 52 74 77 105 111 129 132
Conc manow	Cionicen	101111002	210 317 510-511 545 549 581 598 628
			638 724 766 789 844 860 868 873 919
			927 952 963 968 976 1042 1111 1141
			1160-1161 1229 1266 1346
bone marrow	Clontech	BMD004	111 238 282 549 1083
adult colon	Invitrogen	CLN001	52 260 264 299 494 536 545 564 592
50101			844 873 877 952 976 1042 1152 1268
			1336-1337
adult cervix	BioChain	CVX001	49 51 129 132 151 205 207 238 332-
-aut out viv	Dioonani	012001	335 365-367 392-401 440 466 470-471
			518 537 597 629 832 877 927 976 1006
		1	1085 1117 1129-1134 1192 1202-1205
			1219 1309-1328
diaphragm	BioChain	DIA002	
diaphragm	BioChain	DIA002	74 976 1083

Tissue Origin	RNA Source	Hyseq Library Name	SEQ ID NOS:
endothelial cells	Strategene	EDT001	32 40-41 49 74 79 101 111 120 132
			138 151 204-206 215-217 238 269 316
		}	414 433 505 510 513 550 555 580 582
	<b>\</b>	(	596 675 722 745 798 814 836-841 851
		}	918 976 1041 1043 1073 1083 1131
			1331
Genomic clones	Genomic DNA	EPM001	525-532 927
from the short arm	from Genetic	)	
of chromosome 8	Research		
Genomic clones	Genomic DNA	EPM003	47 525
from the short arm	from Genetic		
of chromosome 8	Research		
Genomic clones	Genomic DNA	EPM004	525 927
from the short arm	from Genetic		
of chromosome 8	Research	TP3 600 f	
Genomic clones	Genomic DNA	EPM005	531
from the short arm of chromosome 8	from Genetic		
	Research	77.0000	74 100 000
esophagus fetal brain	BioChain	ESO002	74 138 238
	Clontech	FBR001	441-442 927
fetal brain	Cloutech	FBR004	215 893 927 1001
fetal brain	Clontech	FBR006	48 61 101 120 132 138 140 147 208
	}	Į	225 271 317 319 336 359 368 405-414
			519 550 571 594 686 715 722 764 824
	Ì	ĺ	829 836 859 909 927 943 947 963 1057
:	1	)	1067-1068 1104 1135-1140 1162 1206-
			1207 1235 1268 1288 1307-1308 1319 1338-1350
fetal brain	Clontech	FBRs03	111 446
fetal brain	Invitrogen	FBT002	41 51 120 151 192-194 264 504 512
letai biani	invinogen	FB1002	535 683 761 798 820-827 844 876 909
	j		963 976 1026 1048 1083 1144 1302
fetal heart	Invitrogen	FHR001	446 566 761
fetal kidney	Clontech	FKD001	51 74 111 127 140 151 184 294 537
	O.O.A.COM	, 1125001	550 630-631 1319
fetal kidney	Clontech	FKD002	111 976 1083
fetal kidney	Invitrogen	FKD007	238 974
fetal lung	Clontech	FLG001	463 566 976 1074 1083 1093
fetal lung	Invitrogen	FLG003	41 238 330 407 415-416 537 573 844
			859 1048 1083 1116 1192
fetal liver-spleen	Columbia	FLS001	8 14 34-35 37 41 43 49 51 54-56 63-64
	University		69-71 74 77 79 87-90 101 107 110-111
			114 120 128-131 138 140 147 150-155
			197 210 215 217 225 238 312 367 384
			414 440 446 460 468 483 496 504-507
			511-515 518-519 523 533-535 537 541
			544-545 547-550 555-560 564 566 571
			577 582 585-586 598 636 646-647 649
			652 664 698 709-710 714 722-723 731
	}	,	735-736 746-753 761 784 798 823 829
		•	832 844 851 858-859 868 873 876 898
			927 943 949 952 963 976 984 1002
			1021 1023 1040 1042 1044 1050 1083
			1093 1116 1120 1129 1131 1144 1174
	}		1217 1251 1254 1256 1302 1308 1311
fatal liver calesa	Columbia	ET SOOO	1319
fetal liver-spleen	Columbia	FLS002	8 36-37 41-46 49 54 64 71 74 79 101
	University		111 120 129 147 207 210 215-216 238 250 330 353 359 366 383-384 414 478
			505 508-509 511 515-524 534-535 537
			544-545 564 566 571 577 591 598 638
	_l	<u> </u>	31 F 3 T 3 3 O T 3 O O 3 I 1 3 I 1 3 7 0 3 0 3 0 3 0 3 0 3 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 0 3 0 3 0 0 0 3 0 0 0 3 0 0 0 3 0 0 0 3 0 0 0 0 3 0 0 0 0 3 0

Tissue Origin	RNA Source	Hyseq Library Name	SEQ ID NOS:
			663 671 698 714 722 725 727 751 798 851 859 873 876 909 927 949 952 983- 984 1002 1023 1042-1044 1085 1095 1131 1144 1178 1199 1233 1240-1270 1331 1340
fetal liver-spleen	Columbia University	FLS003	64 535 976 1256
fetal liver	Invitrogen	FLV001	8 101 120 138 217 446 468 535 566 580 722 730 749 844 918 943 976 1051 1256 1331
fetal liver	Clontech	FLV004	537 926 1256
fetal muscle	Invitrogen	FMS001	51 111 264 312 369-370 404 417-421 425 535 537 577 598 614 836 857 1141 1208 1268
fetal muscle	Invitrogen	FMS002	537
fetal skin	Invitrogen	FSK001	13-26 32 41 51 89 107 111 147 151 225 264 316 405 422-429 488-494 496 519 534-535 537 566 675 732 859 876- 877 898 947 949-950 963 976 1001 1062 1076 1083 1117 1144 1165 1268 1281
fetal skin	Invitrogen	FSK002	537 812
fetal spleen	BioChain	FSP001	87 549
umbilical cord	BioChain	FUC001	27-33 41 49 151 215 238 248-249 301 316 446 495-503 519 521 534-535 537 582 634 691 877 883 927 944-950 963 976 1001 1075 1142-1143 1171 1218 1243 1308
fetal brain	GIBCO	HFB001	41 49 57 79 87 103 111 120 132-135 138 145 151 188 197 207 215 238 264 271 294 316 367 414 440 446 466 504 513-514 535 542-543 550 564 571 596 635 648-654 675 711-715 722-723 798 832 872 876 883 927 976 1095 1144 1168 1171 1178 1211 1335
macrophage	Invitrogen	HMP001	238
infant brain	Columbia University	IB2002	49-50 77 81 89 105 111 136-138 140 151 161 175-179 185 216-217 264 295 299 308-310 371-373 462 476 504 511- 513 533 537 564 566 571 655-657 662 683 716-720 723 752 790-803 829 832 858-859 876 898 909 949 976 1045- 1047 1076-1087 1090 1093 1116 1122 1144 1209-1213 1225 1233 1256 1319 1341
infant brain	Columbia University	IB2003	41 50 77 104 132 215 238 508 512-513 519 566 655 714 794 918 943 976 1067 1092-1093 1233
infant brain	Columbia University	IBM002	311 472-473 753 1214
infant brain	Columbia University	IBS001	51 111 376 474 790 876 949 1144 1204 1221
lung, fibroblast	Strategene	LFB001	151 316 462 514 534 582 675 939 1131
lung tumor	Invitrogen	LGT002	1-7 41 74 79 94 115 120 138-139 156 215 217 269 280 296 337 374-375 384 404 446 454 475-480 498 514 518-519 522 537 545 564 577 597 653 658 705 721-724 754-756 779 859 868 872-874 876-877 919 927 949 951-952 959 976 1002 1042 1048-1053 1076 1083 1088- 1089 1131 1144-1147 1216-1218 1229

Tissue Origin	RNA Source	Hyseq Library Name	SEQ ID NOS:
			1293 1311
lymphocytes	ATCC	LPC001	41 74 111 132 151 253 316 446 550
			634 844 927 976 1085 1268
leukocyte	GIBCO	LUC001	8 11 41 74 86 91-98 101 109 111 120
			147 151 212 215 218 238 252 288 312-
			314.316 338 359 408 427 443-447 505
			510 512 514 518 534 545 549-550 561
	1		564 566 571 577 580 582 587-609 615
			632-638 658-659 698 714 725-728 832
	j		836 841 859 866 873-874 882-883 918-
	!		919 927 943 952 963 976 1042 1076
			1083 1090 1148 1152 1168 1195 1219-
			1220 1224
leukocyte	Clontech	LUC003	74 100 215 232 238 339-341 446 545
	j		657 660 729 873 883 927 952 963 1008
			1042 1116 1120 1149-1150 1215 1222
Melanoma from cell	Clontech	MEL004	210 215 238 342 534 545 592 722 873
line ATCC #CRL			919 929 939 952 976 1071 1118 1218
1424			1235 1245
mammary gland	Invitrogen	MMG001	8-10 40-41 49 73 80 114 138-140 147
	(		217 250-256 264 297-299 305 377-378
	1		398 446 481-486 505 512 537 545 549
		•	571 592 725 730-733 816 829 836 844
	Ì		868 873 876-877 898 926 943 951-960
			963 976 995 1034 1042 1048 1054-
	1		1055 1076 1083 1091 1093 1116-1117
			1124 1152 1302
induced neuron cells	Strategene	NTD001	39 101 111 138 238 361 1225 1251
•			1319
retinoid acid induced	Strategene	NTR001	74 225 976
neuronal cells			
neuronal cells	Strategene	NTU001	129 225 238 304 313 361 657 976
pituitary gland	Clontech	PIT004	976
placenta	Clontech	PLA003	38 976
prostate	Clontech	PRT001	111 188 238 257-258 564 724 961-966
			1067 1095
rectum	Invitrogen	REC001	238 430-431 841 859 868 963 1001
			1116
salivary gland	Clontech	SAL001	8 151 402 432-433 446 496 868 952
			976 1083 1120 1151 1184
small intestine	Clontech	SIN001	8 101 147 215 259-266 446 462 505
			545 592 660 789 836 866 873 927 952
		1	963 967-978 1042 1120 1152 1223-
			1224
skeletal muscle	Clontech	SKM001	238 302 927 943 992 1031
spinal cord	Clontech	SPC001	74 111 132 151 215-216 238 264 267-
			270 343-344 353 379 516 537 566 740
			828 927 976 979-994 1092 1153-1159
			1225 1250
adult spleen	Clontech	SPLc01	698 859 1042
stomach	Clontech	STO001	210 238 271-272 537 580 705 918 952
			995 1171
thalamus	Clontech	THA002	61 219-220 273-276 312 315 330 596
			963 996-1007 1059 1093 1160-1162
thymus	Clonetech	THM001	8 120 151 208 221 316-317 353 639
			750 867 874 878-881 927 963 1023
	ļ	1	1083 1094-1096 1124
thymus	Clontech	THMc02	8 61 114 129 132 210 225 231 306
,	515115511		317-319 336 340 359 380 398 446 448-
		]	463 512 519 545 554 587 598 698 724-
			725 789 812 836 868 873 927 947 952
	<u> </u>	<u> </u>	120 107 012 000 000 010 721 771 702

Tissue Origin	RNA Source	Hyseq Library Name	SEQ ID NOS:
			976 1007 1042 1083 1085 1097-1116
			1122 1147 1177 1226-1229 1234 1311
			1313
thyroid gland	Clontech	THR001	14 41 49 76 94 111 144 151 183 188
		<b>k</b>	210 217 222 253 264 271 277-286 294
			320-326 345-352 361 381-382 446 467
			483 514 534 549-550 564 578 602 649
			844 882-883 927 950 956 976 1008-
	}	}	1028 1076 1083 1117-1120 1142 1163-
		1	1175 1230-1238 1308
trachea	Clontech	TRC001	223-225 238 287 353-354 514
			545 592 611 873 883-884 927
			952 1029-1031 1042 1151-1152
			1170 1176-1177 1239
uterus	Clontech	UTR001	151 226 288-290 355 537 877
			885-886 976 1001 1032-1033
			1232

# TABLE 2

SEQ	Accession	Species	Description	Smith-	%
ID NO:	No.			Waterman Score	Identity
1	B02829	Homo sapiens	Human G protein coupled receptor hRUP5 protein SEQ ID NO:10.	460	100
2	G03564	Homo sapiens	Human secreted protein, SEQ ID NO: 7645.	111	51
3	R26173	Homo sapiens	Part of Major Yo paraneoplastic antigen (CDR62) encoded by clone pY2.	293	76
4	L29536	Homo sapiens	calcium channel L-type alpha 1 subunit	191	65
5	Y94943	Homo sapiens	Human secreted protein clone yt14_1 protein sequence SEQ ID NO:92.	251	50
6	M11507	Homo sapiens	transferrin receptor	120	95
7	AF099100	Homo sapiens	WD-repeat protein 6	1941	93
8	Y92338	Homo sapiens	Human cancer associated antigen precursor from clone NY-REN-45.	245	82
9	G01343	Homo sapiens	Human secreted protein, SEQ ID NO: 5424.	226	91
10	AJ133798	Homo sapiens	copine VII protein	1127	68
11	G02449	Homo sapiens	Human secreted protein, SEQ ID NO: 6530.	584	99
12	X98330	Homo sapiens	ryanodine receptor 2	282	78
13	AL024498	Homo sapiens	dJ417M14.2 (novel serine/threcnine-protein kinase (ortholog of mouse and rat MAK (male germ cell-associated kinase))	293	100
14	AF045577	Pan troglodytes	olfactory receptor OR93Ch	191	36
15	G03131	Homo sapiens	Human secreted protein, SEQ ID NO: 7212.	93	39
16	U26595	Rattus norvegicus	prostaglandin F2a receptor regulatory protein precursor	569	89
17	B08918	Homo sapiens	Human secreted protein sequence encoded by gene 28 SEQ ID NO:75.	99	44
18	Y36203	Homo sapiens	Human secreted protein #75.	165	75
19	U15647	Mus musculus	reverse transcriptase	106	40
20	G02701	Homo sapiens	Human secreted protein, SEQ ID NO: 6782.	544	100
21	Y35923	Homo sapiens	Extended human secreted protein sequence, SEQ ID NO. 172.	1691	100
22	G04030	Homo sapiens	Human secreted protein, SEQ ID NO: 8111.	380	96
23	G02455	Homo sapiens	Human secreted protein, SEQ ID NO: 6536.	123	50
24	AF036329	Homo sapiens	gonadotropin-releasing hormone precursor, second form	284	90
25	G04067	Homo sapiens	Human secreted protein, SEQ ID NO: 8148.	96	32
26	S80119	Rattus sp.	reverse transcriptase homolog	100	34
27	U83303	Homo sapiens	line-1 reverse transcriptase	101	35
28	G03267	Homo sapiens	Human secreted protein, SEQ ID NO: 7348.	135	45

SEQ	Accession	Species	Description	Smith-	%
ID NO:	No.		-	Waterman Score	Identity
29	G04067	Homo sapiens	Human secreted protein, SEQ ID NO: 8148.	83	42
30	G02872	Homo sapiens	Human secreted protein, SEQ ID NO: 6953.	116	72
31	G03371	Homo sapiens	Human secreted protein, SEQ ID NO: 7452.	96	67
32	G03224	Homo sapiens	Human secreted protein, SEQ ID NO: 7305.	58	32
33	Y66688	Homo sapiens	Membrane-bound protein PRO1152.	2457	98
34	Y87071	Homo sapiens	Human secreted protein sequence SEQ ID	348	95
25	TUESTON	1	NO:110.	102	48
35	U15131	Homo sapiens Homo sapiens	p126	182	90
36	Y73464		Human secreted protein clone yl4_1 protein sequence SEQ ID NO:150.	982	'
37	AL133215	Homo sapiens	bA108L7.6 (semaphorin 4G (sema domain, immunoglobulin domain (Ig), transmembrane domain (TM) and short cytoplasmic domain))	687	99
38	AC067969	amino acids 3338-4088	Homo sapiens ryanodine receptor 1 (skeletal)	386	66
39	AL031588	Homo sapiens	dJ1163J1.1 (mostly supported by GENSCAN, FGENES and GENEWISE)	493	76
40	G03628	Homo sapiens	Human secreted protein, SEQ ID NO: 7709.	110	51
41	AF132969	Homo sapiens	CGI-35 protein	228	68
42	Y36268	Homo sapiens	Human secreted protein encoded by gene 45.	220	88
43	X61048	Hydra sp.	mini-collagen	105	35
44	M76546	Helianthus annuus	hydroxyproline-rich protein	110	31
45	U82288	Caenorhabditi s elegans	Rac-like GTPase	139	70
46	G03477	Homo sapiens	Human secreted protein, SEQ ID NO: 7558.	118	58
47	AF090942	Homo sapiens	PRO0657	113	63
48	G03564	Homo sapiens	Human secreted protein, SEQ ID NO: 7645.	90	59
49	AJ005560	Mus musculus	SPR2B protein	72	56
50	G02450	Homo sapiens	Human secreted protein, SEQ ID NO: 6531.	385	98
51	Y91649	Homo sapiens	Human secreted protein sequence encoded by gene 60 SEQ ID NO:322.	973	94
52	U93563	Homo sapiens	putative p150	105	38
53	Y55927	Homo sapiens	Human STLK2 protein.	699	85
54	G02607	Homo sapiens	Human secreted protein, SEQ ID NO: 6688.	145	56
55	AB008175	Mus musculus	hepatic nuclear factor 1-beta short form	356	74
56	M68941	Homo sapiens	protein-tyrosine phophatase	165	41
57	AL031600	Homo sapiens	c390E6.1 (chloride channel 7)	338	76
58	AF011417	Mus	putative pheromone receptor	143	55
59	AF167320	Mus Mus	zinc finger protein ZFP113	558	68
	1772026	musculus		262	06
60	U73036	Homo sapiens	interferon regultory factor 7	263	96 69
61	X07984	Mus musculus	protein-tyrosine kinase	297	09
62	Y29861	Homo sapiens	Human secreted protein clone cb98_4.	791	98
63	U35376	Homo sapiens	repressor transcriptional factor	485	65
64	AF265555	Homo sapiens	ubiquitin-conjugating BIR-domain enzyme APOLLON	785	74
65	G03883	Homo sapiens	Human secreted protein, SEQ ID NO: 7964.	88	95
66	AF177390	Manduca sexta	antennal specific membrane protein AMP	274	54
67	AB040800	Homo sapiens	SREB2	614	100
68	AF030027	Equine herpesvirus 4	24	213	26
69	G02965	Homo sapiens	Human secreted protein, SEQ ID NO: 7046.	261	95
70	W75770	Homo sapiens	Human oxidoreductase YTFO3.	1144	98
71	AB011135	Homo sapiens	KIAA0563 protein	239	76
72	AB011133	Halocynthia	HrPOPK-1	813	78
		roretzi			73
73	AF045454	Cavia porcellus	phospholipase B	955	
74	J02870	Mus	laminin receptor ·	308	61

ID NO: 75 76 77 78 79 80 81 82 1	Accession No. Y00826 AF117754 Y38422 Y14596 Y14591	musculus Rattus norvegicus Homo sapiens Homo sapiens	gp210 (AA 1-1886)  thyroid hormone receptor-associated protein complex component TRAP240	Smith- Waterman Score	Identity 84
75 76 77 78 79 80 79 81 79 82 1	AF117754 Y38422 Y14596	Rattus norvegicus Homo sapiens	thyroid hormone receptor-associated protein	413	84
75 76 77 78 79 80 79 81 79 82 1	AF117754 Y38422 Y14596	Rattus norvegicus Homo sapiens	thyroid hormone receptor-associated protein		
76	AF117754 Y38422 Y14596	Rattus norvegicus Homo sapiens	thyroid hormone receptor-associated protein		
76	AF117754 Y38422 Y14596	norvegicus Homo sapiens Homo sapiens	thyroid hormone receptor-associated protein		
77 78 79 80 81 82	Y38422 Y14596	Homo sapiens Homo sapiens		351	<del></del>
78 79 80 81 82	Y14596	Homo sapiens			54
78 79 80 81 82	Y14596				
78 79 80 81 82	Y14596		Human secreted protein.	468	76
79 80 81 82		and and burgares	Human T-type voltage-gated Ca channel alpha-	1357	99
80 81 82	Y14591		1-I (hCavT3).	1337	
80 81 82		Human	APM-1 protein	767	100
81		papillomaviru	- A protein	] '`'	
81		s type 68		}	Ì
81	AL137802	Homo sapiens	dJ798A10.2 (KIAA0445 protein)	71	34
82	AP000383	Arabidopsis	protein arginine N-methyltransferase-like protein	359	65
	14 000505	thaliana	proton arginato it month transferato ince protoni	337	05
	L46815	Mus	DNA binding protein Rc	895	75
02	240012	musculus	Divi small ploten Re	0,5	1,5
733 I f	G01600	Homo sapiens	Human secreted protein, SEQ ID NO: 5681.	315	96
	Y53886	Homo sapiens	A suppressor of cytokine signalling protein	538	71
~	10000	Troute suprens	designated HSCOP-6.	330	'`
85	AB029002	Homo sapiens	KIAA1079 protein	134	42
	Y28678	Homo sapiens	Human cw272 7 secreted protein.	325	62
	Y99368	Homo sapiens	Human PRO1326 (UNQ686) amino acid	156	48
0/	1 99308	Homo sapiens	sequence SEQ ID NO:100.	136	46
88	AJ225124	Mus	hyperpolarization-activated cation channel,	487	95
00	AJ223124	musculus	HAC3	467	93
89	AE177002		cerebral cell adhesion molecule	200	
	AF177203	Homo sapiens	1	290	56
	Y28280	Homo sapiens	Human G-protein coupled receptor GRIR-2.	326	79
	L39891	Homo sapiens	polycystic kidney disease-associated protein	1751	95
	AF064876	Homo sapiens	ion channel BCNG-1	953	99
	AF170723	Homo sapiens	protein kinase STK10	401	53
94	X13292	Trypanosoma	GPI-phospholipase C (AA 1 - 358)	151	37
		brucei			
	Y34127	Homo sapiens	Human potassium channel K+Hnov11.	661	99
96	X03638	Rattus	sodium channel protein I (aa 1-2009)	1775	92
		norvegicus			<u> </u>
	AF134213	Homo sapiens	ubiquitin-specific protease	1995	99
	G00838	Homo sapiens	Human secreted protein, SEQ ID NO: 4919.	213	38
99	AF021935	Rattus	mytonic dystrophy kinase-related Cdc42-binding	675	48
		norvegicus	kinase		
	AF279265	Homo sapiens	putative anion transporter 1	867	98
101	AC007878	Homo sapiens	match to nuclear protein, NP220; note: sequence	160	60
			difference at residue 58		
102	U22829	Mus	P2Y purinoceptor	264	42
		musculus			
103	Y45023	Homo sapiens	Human sensory transduction G-protein coupled	516	99
	· · · · · · · · · · · · · · · · · · ·		receptor-B3.		
	Y94990	Homo sapiens	Human secreted protein vb21_1, SEQ ID NO:20.	787	98
105	Y87342	Homo sapiens	Human signal peptide containing protein HSPP-	343	57
			119 SEQ ID NO:119.		
	AF169312	Homo sapiens	hepatic angiopoietin-related protein	212	67
	AF116657	Homo sapiens	PRO1310	74	52
108	AE000401	Escherichia	sialic acid transporter	587	96
		coli			1
109	Y38395	Homo sapiens	Human secreted protein encoded by gene No. 10.	693	100
	Y78801	Homo sapiens	Hydrophobic domain containing protein clone	182	94
			HP00631 amino acid sequence.		
111 2	Z25535	Homo sapiens	nuclear pore complex protein hnup153	464	85
	Y94939	Homo sapiens	Human secreted protein clone ye90_1 protein	274	51
	_		sequence SEQ ID NO:84.		1
	AF016365	Homo sapiens	hexokinase 1 isoform td	301	71
	AC007956	Homo sapiens	unknown	520	75
113	ふしいけんさいし	I TOTAL PRINTED I			
113 A		Homo saniens	protein-tyrosine phosphatase	251	1 97
113 1 114 1 115 1	M83738	Homo sapiens	protein-tyrosine phosphatase	251 484	92
113 1 114 1 115 1		Homo sapiens Homo sapiens	protein-tyrosine phosphatase dJ875K15.1.1 (ets homologous factor (ets- domain transcription factor ESE-3A, isoform 1))	251 484	91

December   No.	SEQ	Accession	Species	Description	Smith-	%
190		No.				Identity
190	118	L41816	Homo sapiens	cam kinase I	407	62
AF026954   Bos taurus			Rattus	phosphatidylinositol 3-kinase	627	93
122   123   124   124   125   124   125   124   125   124   125   124   125   125   125   126   126   125   125   126	120	AF026954		subunit precursor; PDPr	1646	94
124   V84107	121	S39392	Homo sapiens		373	68
124				oncostatin-M specific receptor beta subunit	262	
Selegans						
AB021861				•		29
My0360	125		Homo sapiens	guanine nucleotide binding protein beta subunit 4	693	90
128	126	AB021861		apoptosis signal-regulating kinase 2	153	65
129   D32202   Homo sapiens   alpha IC adrenergic receptor isoform 2   574   86   130   AF208043   Homo sapiens   IF116b   496   67   67   67   67   67   67   67	127	AF305210	Homo sapiens		807	97
130						
131						
			Homo sapiens	IFI16b		
133		AF201734	1	testis specific serine kinase-3	800	87
134   W74802   Homo sapiens   Human secreted protein encoded by gene 73   1157   87	132	AF112886	Bos taurus		159	74
Colone HSQEL_25.   Colone HSQEL_25.	133	AJ278314	Homo sapiens	phospholipase C-beta-1b	554	85
136   W80408   Homo sapiens   A secreted protein encoded by clone dt674_2.   866   98     137	134	W74802		clone HSQEL25.	1157	87
137			Homo sapiens			
Similarity to P49205 (PID:g1345860)	136				866	98
AB024034	137	AC002563	1	putative RHO/RAC effector protein; 95% similarity to P49205 (PID:g1345860)	5041	99
thaliana	138	Y96736	Homo sapiens	PRO3434, a novel secreted protein.		100
141	139	AB024034	thaliana		147	55
142	140	W97809		Human GTPase regulator GRAF.	248	56
143   W26642   Homo sapiens   Human RECK cancer-inhibiting protein.   641   82     144		1		Human PLA2 protein.	125	46
144			norvegicus		623	93
145   AF264014   Homo sapiens   Scavenger receptor cysteine-rich type 1 protein   M160 precursor   M160 pr				Human RECK cancer-inhibiting protein.		82
M160 precursor   M160				transmembrane receptor UNC5H2	578	84
147         M96264         Homo sapiens         galactose-1-phosphate uridyl transferase         513         81           148         D64014         Escherichia coli         HrsA         818         90           149         M83316         Escherichia coli         pppGpp phosphohydrolase coli         915         95           150         AL163279         Homo sapiens homolog to cAMP response element binding and beta transducin family proteins         1261         99           151         AF179867         Homo sapiens STE20-like kinase         940         99           152         R95332         Homo sapiens Tumor necrosis factor receptor 1 death domain ligand (clone 3TW).         392         61           153         AF151859         Homo sapiens CGI-101 protein         370         92           154         X66957         Homo sapiens alternatively spliced form         489         81           155         Y16355         Homo sapiens Human secreted protein, SEQ ID NO: 4938.         349         78           157         AF159455         Mus musculus         zinc finger protein         352         74           158         L76191         Homo sapiens         interleukin-1 receptor-associated kinase         537         76           159         AP001743         Homo sap				M160 precursor	727	92
148         D64014         Escherichia coli         HrsA         818         90           149         M83316         Escherichia coli         pppGpp phosphohydrolase         915         95           150         AL163279         Homo sapiens         homolog to cAMP response element binding and beta transducin family proteins         1261         99           151         AF179867         Homo sapiens         STE20-like kinase         940         99           152         R95332         Homo sapiens         Tumor necrosis factor receptor 1 death domain ligand (clone 3TW).         392         61           153         AF151859         Homo sapiens         CGI-101 protein         370         92           154         X66957         Homo sapiens         hexokinase type 1         489         81           155         Y16355         Homo sapiens         alternatively spliced form         432         92           156         G00857         Homo sapiens         Human secreted protein, SEQ ID NO: 4938.         349         78           157         AF159455         Mus musculus         zinc finger protein         352         74           158         L76191         Homo sapiens         interleukin-1 receptor-associated kinase         537         76	146	W63683	Homo sapiens	Human secreted protein 3.	140	40
Coli			Homo sapiens	galactose-1-phosphate uridyl transferase		81
Coli		D64014		HrsA	818	
beta transducin family proteins   151			coli		j	
151         AF179867         Homo sapiens         STE20-like kinase         940         99           152         R95332         Homo sapiens         Tumor necrosis factor receptor 1 death domain ligand (clone 3TW).         392         61           153         AF151859         Homo sapiens         CGI-101 protein         370         92           154         X66957         Homo sapiens         hexokinase type 1         489         81           155         Y16355         Homo sapiens         alternatively spliced form         432         92           156         G00857         Homo sapiens         Human secreted protein, SEQ ID NO: 4938.         349         78           157         AF159455         Mus musculus         zinc finger protein         352         74           158         L76191         Homo sapiens         interleukin-1 receptor-associated kinase         537         76           159         AP001743         Homo sapiens         putative gene, ankirin like, possible dual specifity Ser/Thr/Tyr kinase domain         670         98           160         AJ250425         Rattus norvegicus         Collybistin I         556         74	150	AL163279	Homo sapiens		1261	99
152   R95332   Homo sapiens   Tumor necrosis factor receptor 1 death domain ligand (clone 3TW).   153   AF151859   Homo sapiens   CGI-101 protein   370   92   154   X66957   Homo sapiens   hexokinase type 1   489   81   155   Y16355   Homo sapiens   alternatively spliced form   432   92   156   G00857   Homo sapiens   Human secreted protein, SEQ ID NO: 4938.   349   78   157   AF159455   Mus   zinc finger protein   352   74   158   L76191   Homo sapiens   interleukin-1 receptor-associated kinase   537   76   159   AP001743   Homo sapiens   putative gene, ankirin like, possible dual   670   98   specifity Ser/Thr/Tyr kinase domain   160   AJ250425   Rattus   Collybistin I   556   74	151	AF179867	Homo sapiens		940	99
153         AF151859         Homo sapiens         CGI-101 protein         370         92           154         X66957         Homo sapiens         hexokinase type 1         489         81           155         Y16355         Homo sapiens         alternatively spliced form         432         92           156         G00857         Homo sapiens         Human secreted protein, SEQ ID NO: 4938.         349         78           157         AF159455         Mus musculus         zinc finger protein         352         74           158         L76191         Homo sapiens         interleukin-1 receptor-associated kinase         537         76           159         AP001743         Homo sapiens         putative gene, ankirin like, possible dual specifity Ser/Thr/Tyr kinase domain         670         98           160         AJ250425         Rattus norvegicus         Collybistin I         556         74	152	R95332		Tumor necrosis factor receptor 1 death domain	392	61
154         X66957         Homo sapiens         hexokinase type 1         489         81           155         Y16355         Homo sapiens         alternatively spliced form         432         92           156         G00857         Homo sapiens         Human secreted protein, SEQ ID NO: 4938.         349         78           157         AF159455         Mus musculus         zinc finger protein         352         74           158         L76191         Homo sapiens         interleukin-1 receptor-associated kinase         537         76           159         AP001743         Homo sapiens         putative gene, ankirin like, possible dual specifity Ser/Thr/Tyr kinase domain         670         98           160         AJ250425         Rattus norvegicus         Collybistin I         556         74	153	AF151859	Homo sapiens		370	92
155         Y16355         Homo sapiens         alternatively spliced form         432         92           156         G00857         Homo sapiens         Human secreted protein, SEQ ID NO: 4938.         349         78           157         AF159455         Mus musculus         zinc finger protein         352         74           158         L76191         Homo sapiens         interleukin-1 receptor-associated kinase         537         76           159         AP001743         Homo sapiens         putative gene, ankirin like, possible dual specifity Ser/Thr/Tyr kinase domain         670         98           160         AJ250425         Rattus norvegicus         Collybistin I         556         74						
156         G00857         Homo sapiens         Human secreted protein, SEQ ID NO: 4938.         349         78           157         AF159455         Mus musculus         zinc finger protein         352         74           158         L76191         Homo sapiens interleukin-1 receptor-associated kinase         537         76           159         AP001743         Homo sapiens putative gene, ankirin like, possible dual specifity Ser/Thr/Tyr kinase domain         670         98           160         AJ250425         Rattus norvegicus         Collybistin I         556         74		Y16355		alternatively spliced form	432	
AF159455 Mus musculus  158 L76191 Homo sapiens interleukin-1 receptor-associated kinase 537 76  159 AP001743 Homo sapiens putative gene, ankirin like, possible dual specifity Ser/Thr/Tyr kinase domain  160 AJ250425 Rattus norvegicus  Collybistin I 556 74				Human secreted protein, SEQ ID NO: 4938.	349	78
AP001743 Homo sapiens putative gene, ankirin like, possible dual specifity Ser/Thr/Tyr kinase domain  Rattus norvegicus Collybistin I 556 74			Mus musculus			74
AP001743 Homo sapiens putative gene, ankirin like, possible dual specifity Ser/Thr/Tyr kinase domain  Rattus norvegicus Collybistin I 556 74	158	L76191	Homo sapiens	interleukin-1 receptor-associated kinase	537	76
160 AJ250425 Rattus Collybistin I 556 74	159	AP001743		putative gene, ankirin like, possible dual	670	98
	160				556	74
	161	G02885		Human secreted protein, SEQ ID NO: 6966.	370	100

SEQ ID	Accession No.	Species	Description	Smith- Waterman	% Identity
NO:	1			Score	
162	Z22968	Homo sapiens	M130 antigen	610	100
163	AF181121	Homo sapiens	ATP-dependent Ca2+ pump PMR1	336	92
164	AF055636	Homo sapiens	leucine-rich glioma-inactivated protein precursor	455	94
165	AF160798	Rattus norvegicus	calcium transporter CaT1	700	96
166	Y76332	Homo sapiens	Fragment of human secreted protein encoded by gene 38.	327	45
167	Y48607	Homo sapiens	Human breast tumour-associated protein 68.	1072	99
168	AB020741	Mus	NIK-related kinase	197	43
169	AF252293	musculus Homo sapiens	PAR3	596	44
170	U59429	Cricetinae	diacylglycerol kinase eta	481	82
		gen. sp.			
171	AF035268	Homo sapiens	phosphatidylserine-specific phospholipase A1	386	42
172	AF127085	Mus musculus	semaphorin cytoplasmic domain-associated protein 3B	507	82
173	Y27918	Homo sapiens	Human secreted protein encoded by gene No. 123.	653	99
174	G02979	Homo sapiens	Human secreted protein, SEQ ID NO: 7060.	538	97
175	U36488	Mus musculus	embryonic stem cell phosphatase	168	55
176	W95629	Homo sapiens	Homo sapiens secreted protein gene clone gm196 4.	1022	100
177	AF289023	Homo sapiens	formiminotransferase cyclodeaminase form D	255	93
178	X04936	Homo sapiens	T-cell receptor alpha-chain (413 is 2nd base in	710	99
			codon)		
179	AF127481	Homo sapiens	non-ocogenic Rho GTPase-specific GTP exchange factor	175	80
180	G00978	Homo sapiens	Human secreted protein, SEQ ID NO: 5059.	517	94
181	Y66645	Homo sapiens	Membrane-bound protein PRO1310.	671	96
182	AF110640	Homo sapiens	orphan seven-transmembrane receptor	862	100
183	AB020854	Bos taurus	orphan transporter short splicing variant	766	84
184	AF169691	Homo sapiens	cadherin-like protein VR8	375	38
185	AF126372	Homo sapiens	thyrotropin-releasing hormone degrading ectoenzyme	985	99
186	L20966	Homo sapiens	phosphodiesterase	541	76
187	G02920	Homo sapiens	Human secreted protein, SEQ ID NO: 7001.	254	93
188	Y94918	Homo sapiens	Human secreted protein clone dd504_18 protein sequence SEQ ID NO:42.	301	98
189	Y66713	Homo sapiens	Membrane-bound protein PRO1309.	694	100
190	G03244	Homo sapiens	Human secreted protein, SEQ ID NO: 7325.	331	73
191	U36771	Rattus norvegicus	sn-glycerol 3-phosphate acyltransferase	707	92
192	R05935	Homo sapiens	Secreted GPIIb subunit of multiple subunit polypeptide (MSP)GPIIb-IIIa.	157	72
193	M92084	Theileria	cascin kinase II alpha subunit	364	50
104	V66645	Homo sapiens	Membrane hound protein PPO1210	448	90
194 195	Y66645 W95631	Homo sapiens Homo sapiens	Membrane-bound protein PRO1310.  Homo sapiens secreted protein gene clone	382	49
196	AF255614	Rattus	hj968_2. scaffolding protein SLIPR	680	99
197	AC021640	norvegicus Arabidopsis	putative phosphatidate phosphohydrolase	300	41
198	AF073967	thaliana Mus	olfactory receptor	316	43
		musculus domesticus			
	W01730	Homo sapiens	Human G-protein receptor HPRAJ70.	617	98
199		Homo sapiens	pancreas-enriched phospholipase C	625	89
	AFIL/948		CDC42-binding protein kinase beta	636	94
200	AF117948 AF128625	Homo caniere			1 77
201	AF128625	Homo sapiens			100
200 201 202		Homo sapiens Homo sapiens Homo sapiens	Link guanine nucleotide exchange factor II Human secreted protein clone qc646_1 protein	1303 701	100 99
199 200 201 202 203 204	AF128625 AF117946	Homo sapiens	Link guanine nucleotide exchange factor II	1303	

SEQ	Accession	Species	Description	Smith-	1%
ID NO:	No.			Waterman Score	Identity
110.	<del></del>	-	{ovarian cancer critical region of deletion}		<u> </u>
206	U18315	Sus scrofa	parathyroid receptor	122	60
207	AF255342	Homo sapiens	putative pheromone receptor V1RL1 long form	170	96
208	S52051	Rattus sp.	neurotransmitter transporter	715	94
209	W63683	Homo sapiens	Human secreted protein 3.	840	99
210	D79992	Homo sapiens	similar to Drosophila photoreceptor cell-specific protein, calphotin.	541	82
211	AF117948	Homo sapiens	pancreas-enriched phospholipase C	1348	99
212	U81035	Rattus norvegicus	ankyrin binding cell adhesion molecule neurofascin	471	69
213	AF154846	Homo sapiens	zinc finger protein	798	56
214	AF102777	Mus musculus	FYVE finger-containing phosphoinositide kinase	933	93
215	AL163303	Homo sapiens	putative gene containing transmembrane domain	523	89
216	U26595	Rattus norvegicus	prostaglandin F2a receptor regulatory protein precursor	563	78
217	G04095	Homo sapiens	Human secreted protein, SEQ ID NO: 8176.	644	98
218	X75756	Homo sapiens	protein kinase C mu	314	81
219	Y66723	Homo sapiens	Membrane-bound protein PRO1100.	770	98
220	D88577	Mus musculus	Kupffer cell receptor	567	40
221	AF258465	Homo sapiens	OTRPC4	853	100
222	AF021935	Rattus norvegicus	mytonic dystrophy kinase-related Cdc42-binding kinase	636	96
223	AL136527	Homo sapiens	bA215B13.1 (A kinase (PRKA) anchor protein 11)	693	100
224	AB032417	Homo sapiens	WNT receptor Frizzled-4	690	99
225	AF030430	Mus musculus	semaphorin VIa	703	68
226	AE000218	Escherichia coli	putative dihydroxyacetone kinase (EC 2.7.1.2)	297	39
227	AF302150	Homo sapiens	phosphoinositol 3-phosphate-binding protein-2	2080	100
228	AB024573	Mus musculus	GTP-binding like protein 2	265	88
229	AF122924	Xenopus laevis	Wnt inhibitory factor-1	316	40
230	G03205	Homo sapiens	Human secreted protein, SEQ ID NO: 7286.	229	100
231	X98260	Homo sapiens	M-phase phosphoprotein 11	265	92
232	R92754	Homo sapiens	Human growth differentiation factor-12.	682	95
233	R75111	Homo sapiens	Glycosyl-phosphatidylinositol-specific phospholipase-D.	290	100
234	W69431	Homo sapiens	Human secreted protein cw1233_3.	235	97
235	Y08686	Homo sapiens	serine palmitoyltransferase, subunit II	859	81
236 237	AF118275 X81466	Homo sapiens Mus	atrophin-related protein ARP Embryo Brain Kinase	117 460	37 62
238	U64857	musculus Caenorhabditi s elegans	similar to the BPTI/Kunitz family of inhibitors; most similar to tissue factor pathway inhibitor precursor (TFPI)	284	33
239	AJ250840	Mus musculus	serine/threonine protein kinase	739	63
240	AJ223472	Mus musculus	transcription elongation factor TFIIS.h	222	38
241	Y94906	Homo sapiens	Human secreted protein clone rb649_3 protein sequence SEQ ID NO:18.	353	52
242	AF169301	Homo sapiens	Na+/sulfate cotransporter SUT-1	591	99
243	L22022	Rattus norvegicus	orphan transporter v7-3	667	93
244	AF016191	Rattus norvegicus	potassium channel	1043	98
245	AF097366	Homo sapiens	cone sodium-calcium potassium exchanger	645	98
246	Y29868	Homo sapiens	Human secreted protein clone pp325 9.	497	98
247	AF180475	Homo sapiens	Not4-Np	188	83
248	Y17227	Homo sapiens	Human secreted protein (clone ya1-1).	690	99
249	AF250910	Manduca	death-associated small cytoplasmic leucine-rich	182	31

SEQ	Accession	Species	Description	Smith-	%
ID	No.	Į		Waterman	Identity
NO:				Score	
0.70	AB100776	sexta	protein SCLP	104	<del>                                     </del>
250	AF192756	Kaposi's	Orf73	134	34
		sarcoma- associated	1	1	ŀ
					l.
051	AD000004	herpesvirus	l Morr	1 200	<del> </del>
251	AB022694	Homo sapiens	MOK protein kinase	209	83
252	W55045	Homo sapiens	Neural adhesion molecule (ethb0018f2 product).	469	100
253	L46815	Mus	DNA binding protein Rc	251	67
254	W68505	musculus		100	<del> </del>
254 255		Homo sapiens	Human acid sensing ionic channel.	173	82
255	AF070066	Mus musculus	Citron-K kinase	1201	98
256	G02491	Homo sapiens	Human secreted protein, SEQ ID NO: 6572.	460	100
257	Z12841	Oryctolagus	Phospholipase	368	80
237	212041	cuniculus	Phospholipase	308	80
258	Y95436	Homo sapiens	Human calcium channel SOC-3/CRAC-2.	1857	99
259	AJ222968	Mus	L-periaxin	430	72
239	AJ222900	musculus	L-periaxiii	430	12
260	AJ250839	Homo sapiens	serine/threonine protein kinase	861	100
261	AJ249977	Homo sapiens	AMP-activated protein kinase gamma 3 subunit	758	98
262	AF141386	Rattus	SLIT-2	198	40
202	AL 141360	norvegicus	0511-2	198	40
263	AF022859	Homo sapiens	neuropilin-2(a0)	335	62
264	AF160477	Homo sapiens	Ig superfamily receptor LNIR precursor	387	91
265	Y44662	Homo sapiens	Human 14273 G-protein coupled receptor	636	99
203	144002	Homo sapiens	(GPCR).	030	99
266	U27269	Mus	sodium glucose cotransporter	204	56
200	02,20)	musculus	Social Election Contains porter	204	1 50
267	AF124491	Homo sapiens	ARF GTPase-activating protein GIT2	159	75
268	AF127389	Rattus	putative taste receptor TR1	209	39
	111 12/303	norvegicus	paramy association and	203	
269	X98296	Homo sapiens	ubiquitin hydrolase	215	95
270	X78482	Streptococcus	Fc-gamma receptor	129	26
		pyogenes		1	
271	AB009883	Nicotiana	KED	109	26
		tabacum			
272	AF137367	Mus	VPS10 domain receptor protein SORCS	899	97
		musculus			
273	L34938	Rattus	ionotropic glutamate receptor	460	86
		norvegicus	<u> </u>		
274	AL022724	Homo sapiens	dJ413H6.1.1 (hamster Androgen-dependent	188	74
			Expressed Protein LIKE PUTATIVE protein)	1	
	17705077	<del> </del>	(isoform 1)	ļ	ļ <u>.</u>
275	AF265555	Homo sapiens	ubiquitin-conjugating BIR-domain enzyme	173	94
000	000000	ļ.,	APOLLON	<del> </del>	
276	G02872	Homo sapiens	Human secreted protein, SEQ ID NO: 6953.	148	56
277	L40380	Homo sapiens	thyroid receptor interactor	430	61
278	AB046851	Homo sapiens	KIAA1631 protein	283	96
279	AC008075	Arabidopsis	Contains PF 00069 Eukaryotic protein kinase	157	43
200	1,000	thaliana	domain.	ļ	<u> </u>
280	M83738	Homo sapiens	protein-tyrosine phosphatase	181	73
281	AK024397	Homo sapiens	unnamed protein product	439	91
282	AF141326	Homo sapiens	RNA helicase HDB/DICE1	497	84
283	AF156530	Mus	ETS-domain transcriptional repressor PE1	605	76
204	170022	musculus		1	1.00
284	Y29336	Homo sapiens	Human secreted protein clone cs756_2 alternate	647	100
205	3/72402	77	reading frame protein.	1 200	100
285	Y73402	Homo sapiens	Human secreted protein clone yc25_1 protein	300	90
206	AP016411	Transaction	sequence SEQ ID NO:26.	127	100
286	AF016411	Homo sapiens	KCNA3.1B	137	100
287	W89253	Homo sapiens	Human ALP.	688	97
288	AF112886	Bos taurus	differentiation enhancing factor 1	750	96
289	AF113131	Homo sapiens	host cell factor homolog LCP	367	44
290	U52111	Homo sapiens	plexin-related protein	698	100
291	AF026504	Rattus	SPA-1 like protein p1294	603	89

SEQ ID	Accession No.	Species	Description	Smith- Waterman	% Identity
NO:				Score	<u> </u>
		norvegicus			
292	AF102854	Rattus norvegicus	membrane-associated guanylate kinase- interacting protein 2 Maguin-2	124	53
293	X99211	Drosophila melanogaster	ubiquitin-specific protease	143	38
294	Y94943	Homo sapiens	Human secreted protein clone yt14_1 protein sequence SEQ ID NO:92.	185	94
295	Y94890	Homo sapiens	Human protein clone HP02798.	108	59
296	AF019767	Homo sapiens	zinc finger protein	154	96
297	Y28568	Homo sapiens	Secreted peptide clone bd577 1.	568	84
298	Y94943	Homo sapiens	Human secreted protein clone yt14_1 protein sequence SEQ ID NO:92.	182	97
299	B08906	Homo sapiens	Human secreted protein sequence encoded by gene 16 SEQ ID NO:63.	605	69
300	R58890	Homo sapiens	Human-32 cadherin-related molecule.	212	97
301	AF022859	Homo sapiens	neuropilin-2(a0)	277	100
302	Y71124	Homo sapiens	Human mitogenic regulator duox2.	716	97
303	Y44297	Homo sapiens	Human receptor tyrosine kinase.	228	97
304	D32050	Homo sapiens	alanyl-tRNA synthetase	192	80
305	U43586	Homo sapiens	protein kinase related to Raf protein kinases; Method: conceptual translation supplied by author	428	72
306	R54872	Homo sapiens	Human H13 viral receptor mutant 4.	280	95
307	D78572	Mus musculus	membrane glycoprotein	199	41
308	AF255614	Rattus norvegicus	scaffolding protein SLIPR	639	88
309	S79463	Mus sp.	semaphorin homolog=M-Sema F	162	89
310	AF178941	Homo sapiens	ATP-binding cassette sub-family A member 2	736	100
311	U03413	Dictyostelium discoideum	calcium binding protein	151	36
312	Y87347	Homo sapiens	Human signal peptide containing protein HSPP- 124 SEQ ID NO:124.	744	100
313	Z97055	Homo sapiens	dJ388M5.4 (putative GS2 like protein)	789 .	99
314	AC004010	Homo sapiens	similar to Leucine-rich transmembrane proteins; 44% similarity to U42767 (PID:g1736918)	197	38
315	AL021392	Homo sapiens	dJ439F8.2 (supported by GENSCAN and GENEWISE)	278	38
316	U70209	Mus musculus	polycystic kidney disease 1 protein	165	38
317	AF109643	Rattus norvegicus	coxsackie-adenovirus-receptor homolog	223	38
318	AF104923	Homo sapiens	putative transcription factor	138	84
319	AF100287	Trypanosoma vivax	activated protein kinase C receptor homolog	141	38
320	G00588	Homo sapiens	Human secreted protein, SEQ ID NO: 4669.	125	51
321	Y21591	Homo sapiens	Human secreted protein (clone CC332-33).	459	97
322	D26070	Homo sapiens	human type 1 inositol 1,4,5-trisphosphate receptor	232	97
323	Y27918	Homo sapiens	Human secreted protein encoded by gene No. 123.	306	88
324	AF010144	Homo sapiens	neuronal thread protein AD7c-NTP	209	70
325	M19650	Homo sapiens	2',3'-cyclic-nucleotide 3'-phosphodiesterase (EC 3.1.4.37)	214	97
326	W80396	Homo sapiens	A secreted protein encoded by clone bp646_10.	140	70
327	X75756	Homo sapiens	protein kinase C mu	540	78
328	G02292	Homo sapiens	Human secreted protein, SEQ ID NO: 6373.	721	99
329	AF168990	Homo sapiens	putative GTP-binding protein	877	99
330	S67984	Homo sapiens	anti-HIV gp120 antibody heavy chain variable region	581	80
331	X13916	Homo sapiens	LDL-receptor related precursor (AA -19 to 4525)	2823	98
332	Y87330	Homo sapiens	Human signal peptide containing protein HSPP- 107 SEQ ID NO:107.	1127	100
333	Y28503	Homo sapiens	HGFH3 Human Growth Factor Homologue 3.	320	98
334	AC002563	Homo sapiens	putative RHO/RAC effector protein; 95%	327	93

SEQ	Accession	Species	Description	Smith-	%
ID	No.			Waterman	Identity
NO:		<u></u>		Score	<u> </u>
			similarity to P49205 (PID:g1345860)		
33 <i>5</i>	Y87347	Homo sapiens	Human signal peptide containing protein HSPP-	1111	67
			124 SEQ ID NO:124.		
336	AF006466	Mus	lymphocyte specific formin related protein	193	75
		musculus			
337	AF265555	Homo sapiens	ubiquitin-conjugating BIR-domain enzyme	632	97
	1		APOLLON		
338	Y13443	Homo sapiens	Amino acid sequence of hSlo3-2.	516	100
339	Y07637	Homo sapiens	putative GABA-gated chloride channel	189	100
340	Y05734	Homo sapiens	Human Grb7 effector 2.2412 protein.	2156	99
341	AE000497	Escherichia	L-idonate transcriptional regulator	928	98
	,	coli		]	
342	D90855	Escherichia	glycerol-3-phosphate dehydrogenase (EC	769	99
	250055	coli	1.1.99.5) chain A, anaerobic	1,05	-
343	D85613	Escherichia	membrane component	399	100
J-1J	D03013	coli	inchiorane component	1 377	100
344	M93239	Escherichia	transmembrane protein	232	100
3 <del>44</del>	10193239		transmemorane protein	232	100
345	M60177	coli Escherichia	onterphentin	759	99
545	MOU1//		enterobactin	139	77
246	- D00000	coli	C	(26	107
346	D90699	Escherichia	Sensor protein copS (EC 2.7.3).	638	97
3.45	Doccus	coli		1.55	1
347	D90843	Escherichia	CapB protein.	552	100
		coli			
348	M13422	Escherichia	49 kd protein	1193	96
		coli			
349	L10328	Escherichia	similar to drug resistance translocases	340	90
		coli			
350	X69942	Mus	enhancer-trap-locus-1	560	82
		musculus	-		
351	AF239613	Homo sapiens	apamin-sensitive small-conductance Ca2+-	463	80
			activated potassium channel	1	1
352	D90777	Escherichia	3-hydroxybutyryl-CoA dehydrogenase (EC	577	100
		coli	1.1.1.157) (b- hydroxybutyryl-CoA	ļ	
			dehydrogenase) (BhbD).		
353	D90863	Escherichia	similar to	311	98
		coli		l	
354	Y52386	Homo sapiens	Human transmembrane protein HP02000.	133	58
355	Y31645	Homo sapiens	Human transport-associated protein-7 (TRANP-	482	55
555	151515	Tromo suprems	7).	1 .02	""
356	Y58637	Homo sapiens	Protein regulating gene expression PRGE-30.	119	51
357	AF119226	Homo sapiens	dual-specificity tyrosine phosphatase YVH1	1788	100
358	Y87219	Homo sapiens	Human secreted protein sequence SEQ ID	165	100
<i>33</i> 6	10/219	1101110 Sapiciis	NO:258.	103	100
359	J00132	IIoma coniona		233	93
		Homo sapiens	beta-fibrinogen		1 -
360	G03789	Homo sapiens	Human secreted protein, SEQ ID NO: 7870.	128	70
361	R28916	Homo sapiens	Type III procollagen (prior art).	108	40
362	U16655	Rattus	phospholipase C delta-4	649	65
	L	norvegicus			
363	G03119	Homo sapiens	Human secreted protein, SEQ ID NO: 7200.	95	42
364	U47276	Gallus gallus	chicken brain factor-2	104	34
365	G03789	Homo sapiens	Human secreted protein, SEQ ID NO: 7870.	183	65
366	G04091	Homo sapiens	Human secreted protein, SEQ ID NO: 8172.	118	46
367	X98258	Homo sapiens	M-phase phosphoprotein 9	564	75
368	AL021366	Homo sapiens	cICK0721Q.3 (Kinesin related protein)	3387	99
369	U70932	Peromyscus	reverse transcriptase	92	59
	1	leucopus			
370	X86400	Homo sapiens	gamma subunit of sodium potassium ATPase	242	73
310	700400	riomo sapiens		242	13
271	002170	YT	like	165	-
371	G03172	Homo sapiens	Human secreted protein, SEQ ID NO: 7253.	165	56
372	U49974	Homo sapiens	mariner transposase	257	55
373	X13916	Homo sapiens	LDL-receptor related precursor (AA -19 to 4525)	21193	99
374	AF234765	Rattus	serine-arginine-rich splicing regulatory protein	1182	78
		norvegicus	SRRP86		
375	U49974	Homo sapiens	mariner transposase	172	55

SEQ	Accession	Species	Description	Smith-	%
ID	No.			Waterman	Identity
NO:	<u> </u>	<del> </del>		Score	<u> </u>
376	G01984	Homo sapiens	Human secreted protein, SEQ ID NO: 6065.	221	67
377	G00669	Homo sapiens	Human secreted protein, SEQ ID NO: 4750.	600	100
378	X52574	Mus musculus	GTP binding protein	1456	91
379	R69095	Homo sapiens	Anti-HIV Fab tat31 light chain.	68	37
380	J04974	Homo sapiens	alpha-2 type XI collagen	125	37
381	AB002405	Homo sapiens	LAK-4p	530	43
382	U64830	Dictyostelium	protein tyrosine kinase	115	44
362	004630	discoideum	protein tyrosine kinase	113	44
383	G02916	Homo sapiens	Human secreted protein, SEQ ID NO: 6997.	618	98
384	G01194	Homo sapiens	Human secreted protein, SEQ ID NO: 5275.	617	93
385	AJ245822	Homo sapiens	type I transmembrane receptor	4560	100
386	D86974	Homo sapiens	KIAA0220	2148	98
387	G03203	Homo sapiens	Human secreted protein, SEQ ID NO: 7284.	142	50
388	G04072	Homo sapiens	Human secreted protein, SEQ ID NO: 8153.	. 99	59
389	M12140	Homo sapiens	envelope protein	197	51
390	AJ293309	Homo sapiens	NHP2 protein	461	77
391	Y42751	Homo sapiens	Human calcium binding protein 2 (CaBP-2).	181	94
392	W48351	Homo sapiens	Human breast cancer related protein BCRB2.	241	66
393	Y14442	Homo sapiens	olfactory receptor protein	339	54
394	W85607	Homo sapiens	Secreted protein clone da228_6.	957	100
395	Y76332	Homo sapiens	Fragment of human secreted protein encoded by gene 38.	171	34
396	G03930	Homo sapiens	Human secreted protein, SEQ ID NO: 8011.	250	100
397	AB032904	Hylobates	dopamine receptor D4	105	35
,,,	1113032301	syndactylus	dopaining receptor D4	105	33
398	AJ007798	Homo sapiens	stromal antigen 3, (STAG3)	861	.85
399	Y91405	Homo sapiens	Human secreted protein sequence encoded by gene 2 SEQ ID NO:126.	1047	92
400	Y29861	Homo sapiens	Human secreted protein clone cb98_4.	162	37
401	D87002	Homo sapiens	similar to rat integral membrane glycoprotein;	527	78
			accession number Z21513.	Ĺ	
402	AF100754	Homo sapiens	ancient ubiquitous protein AUP1 isoform	853	95
403	X74904	Gallus gallus	alpha-2-macroglobulin receptor	258	60
404	AF075462	Mus musculus	ADP-ribosylation factor-directed GTPase activating protein isoform b	545	89
405	X92887	Human endogenous retrovirus K	pol/env	162	30
406	Y30162	Homo sapiens	Human dorsal root receptor 4 hDRR4.	325	72
107	AK022626	Homo sapiens	unnamed protein product	2833	99
408	L13802	Homo sapiens	ribosmal protein small subunit	264	92
409	Y91600	Homo sapiens	Human secreted protein sequence encoded by gene 9 SEQ ID NO:273.	1788	89
410	W88745	Homo sapiens	Secreted protein encoded by gene 30 clone	2004	99
411	AB043953	Mus	HTSEV09. Chat-H	2628	82
		musculus			
412	Y86233	Homo sapiens	Human secreted protein HNTMX29, SEQ ID NO:148.	1014	92
413	U10542	Pan troglodytes	MHC class I A	265	71
414	AF155097	Homo sapiens	NY-REN-7 antigen	850	95
115	G03203	Homo sapiens	Human secreted protein, SEQ ID NO: 7284.	88	48
116	Y57911	Homo sapiens	Human transmembrane protein HTMPN-35.	266	89
17	W27651	Homo sapiens	Secreted protein AT205.	481	60
118	Y76884	Homo sapiens	Retinoblastoma binding protein-7sequence.	3077	87
119	AF255559	Notothenia	alpha tubulin	289	68
		coriiceps	•		_
120	G01984	Homo sapiens	Human secreted protein, SEQ ID NO: 6065.	209	74
121	AL109827	Homo sapiens	dJ309K20.2 (acrosomal protein ACR55 (similar to rat sperm antigen 4 (SPAG4)))	1446	96
422	AC008075	Arabidopsis	F24J5.4	112	35
T44.	WC000013	j zarabiuopsis	1'27JJ.4	112	1 23

SEQ	Accession	Species	Description	Smith-	%
$\mathbf{ID}$	No.			Waterman	Identity
NO:			<u> </u>	Score	
423	AF231705	Homo sapiens	Alu co-repressor 1	1090	100
424	AF234887	Homo sapiens	FLAMINGO 1	6268	97
425	Y35942	Homo sapiens	Extended human secreted protein sequence, SEQ ID NO. 191.	1961	99
426	AB009288	Homo sapiens	N-copine	635	98
427	L12392	Homo sapiens	Huntington's Disease protein	16080	99
428	Y94990	Homo sapiens	Human secreted protein vb21_1, SEQ ID NO:20.	768	98
429	AJ293573	Homo sapiens	zinc finger protein Cezanne	542	87
430	Y84441	Homo sapiens	Amino acid sequence of a human RNA-associated protein.	2074	100
431	G02850	Homo sapiens	Human secreted protein, SEQ ID NO: 6931.	723	95
432	G04067	Homo sapiens	Human secreted protein, SEQ ID NO: 8148.	73	42
433	AF159296	Lycopersicon esculentum	extensin-like protein	613	48
434	W48351	Homo sapiens	Human breast cancer related protein BCRB2.	135	44
435	X73874	Homo sapiens	phosphorylase kinase	3442	97
436	AF161426	Homo sapiens	HSPC308	268	74
437	Y30812	Homo sapiens	Human secreted protein encoded from gene 2.	1055	52
438	G03798	Homo sapiens	Human secreted protein, SEQ ID NO: 7879.	168	56
439	X14766	Homo sapiens	GABA-A receptor alpha 1 subunit	2294	96
440	X02344	Homo sapiens	beta-tubulin	311	95
441	AF168418	Homo sapiens	activating signal cointegrator 1	1882	100
441	L11672		activating signal conficerator 1	795	54
		Homo sapiens	zinc finger protein		1 -
443	G03203	Homo sapiens	Human secreted protein, SEQ ID NO: 7284.	93	26
444	A52140	unidentified	HUMAN NDR	2451	100
445	X98330	Homo sapiens	ryanodine receptor 2	9356	99
446	AF116712	Homo sapiens	PRO2738	227	49
447	AF245447	Homo sapiens	sphingosine kinase type 2 isoform	576	99
448	AF133086	Homo sapiens	membrane-type serine protease 1	2630	94
449	U87305	Rattus norvegicus	transmembrane receptor UNC5H1	817	93
450	AF081249	Homo sapiens	JAW1-related protein MRVI1A long isoform	4568	99
451	AC005498	Homo sapiens	R31665 1	316	62
452	M60235	Homo sapiens	granule membrane protein-140	464	73
453	AB036706	Homo sapiens	intelectin	730	88
454	G00918	Homo sapiens	Human secreted protein, SEQ ID NO: 4999.	263	81
455	Y22634	Homo sapiens	Human cytokine inducible regulatory protein-1 (CIRP-1).	192	67
456	Y36705	Homo sapiens	Fragment of human secreted protein encoded by gene 62.	106	40
457	N91325	Homo sapiens	DNA encoding human growth hormone receptor.	3282	96
458	M19155	Plasmodium	S-antigen precursor	110	36
-		falciparum			
459	Y13377	Homo sapiens	Amino acid sequence of protein PRO257.	509	98
460	Y02693	Homo sapiens	Human secreted protein encoded by gene 44	149	43
461	Y14482	Homo sapiens	clone HTDAD22.  Fragment of human secreted protein encoded by	184	54
462	Y53005	Homo sapiens	gene 17.  Human secreted protein clone pm749_8 protein	135	47
463	X84960	Triticum	sequence SEQ ID NO:16. low molecular weight glutenin	109	33
464	W19919	aestivum  Homo sapiens		1781	85
			Human Ksr-1 (kinase suppressor of Ras).		
465	AF189764	Mus musculus	alpha/beta hydrolase-1	502	59
466	U93569	Homo sapiens	p40	101	30
467	Y41528	Homo sapiens	Fragment of human secreted protein encoded by gene 77.	1172	99
468	G02872	Homo sapiens	Human secreted protein, SEQ ID NO: 6953.	149	52
469	AJ000008	Homo sapiens	PI3-kinase	5832	97
470	X70922	Mus musculus	neurotoxin homologue	118	47
471	G03797	Homo sapiens	Human secreted protein, SEQ ID NO: 7878.	198	75
472	Y36705	Homo sapiens	Fragment of human secreted protein encoded by	72	57

SEQ ID NO:	Accession No.	Species	Description	Smith- Waterman Score	% Identity
			gene 62.		<del> </del>
473	G02313	Homo sapiens	Human secreted protein, SEQ ID NO: 6394.	328	100
474	Y07007	Homo sapiens	Breast cancer associated antigen precursor sequence.	1013	97
475	W93254	Homo sapiens	Human ESRP1 protein.	943	80
476	W48351	Homo sapiens	Human breast cancer related protein BCRB2.	236	65
477	Y02693	Homo sapiens	Human secreted protein encoded by gene 44 clone HTDAD22.	202	60
478	G01870	Homo sapiens	Human secreted protein, SEQ ID NO: 5951.	267	100
479	AF102777	Mus musculus	FYVE finger-containing phosphoinositide kinase	3427	92
480	G03052	Homo sapiens	Human secreted protein, SEQ ID NO: 7133.	123	53
481	W87701	Homo sapiens	A human membrane fusion protein designated SYTAX1.	221	77
482	G03119	Homo sapiens	Human secreted protein, SEQ ID NO: 7200.	131	39
483	AF210651	Homo sapiens	NAG18	124	59
484	AF010144	Homo sapiens	neuronal thread protein AD7c-NTP	343	50
485	G00637	Homo sapiens	Human secreted protein, SEQ ID NO: 4718.	129	70
486	U15174	Homo sapiens	BCL2/adenovirus E1B 19kD-interacting protein 3	149	73
487	Y76167	Homo sapiens	Human secreted protein encoded by gene 44.	627	100
488	AJ275213	Homo sapiens	stabilin-1	1244	91
489	G03798	Homo sapiens	Human secreted protein, SEQ ID NO: 7879.	313	65
490	L12392	Homo sapiens	Huntington's Disease protein	16081	100
491	G03789	Homo sapiens	Human secreted protein, SEQ ID NO: 7870.	197	66
492	J03799	Homo sapiens	laminin-binding protein	228	70
493	U15174	Homo sapiens	BCL2/adenovirus E1B 19kD-interacting protein 3	128	41
494	Y02693	Homo sapiens	Human secreted protein encoded by gene 44 clone HTDAD22.	197	67
495	AC005175	Homo sapiens	R31449_3	889	94
496	G03786	Homo sapiens	Human secreted protein, SEQ ID NO: 7867.	229	61
497	AB030237	Canis familiaris	D4 dopamine receptor	90	48
498	G02872	Homo sapiens	Human secreted protein, SEQ ID NO: 6953.	228	65
499	U70935	Peromyscus maniculatus	reverse transcriptase	213	52
500	U48508	Homo sapiens	skeletal muscle ryanodine receptor	26406	99
501	G03371	Homo sapiens	Human secreted protein, SEQ ID NO: 7452.	105	58
502	AF119851	Homo sapiens	PRO1722	156	62
503	AF113685	Homo sapiens	PRO0974	116	50
504	U79458	Homo sapiens	WW domain binding protein-2	322	59
505	W29651	Homo sapiens	Human secreted protein CD124_3.	608	55
506	W85459	Homo sapiens	Secreted protein encoded by clone dh1135_9.	986	70
507	Y86265	Homo sapiens	Human secreted protein HUSXE77, SEQ ID NO:180.	115	33
508	AL160175	Homo sapiens	bA243J16.3 (similar to MYLK (myosin, light polypeptide kinase))	184	92
509	U43360	Peromyscus maniculatus	reverse transcriptase	97	62
510	G03789	Homo sapiens	Human secreted protein, SEQ ID NO: 7870.	117	63
511	W79092	Homo sapiens	Human secreted protein dn740_3.	1058	100
512	AF010144	Homo sapiens	neuronal thread protein AD7c-NTP	205	64
513	AJ133439	Homo sapiens	GRIP1 protein	2151	100
514	AE003456	Drosophila melanogaster	CG6393 gene product	259	42 .
515	Z17206	Xenopus laevis	p46XIEg22	128	40
516	AF104413	Homo sapiens	large tumor suppressor 1	1766	94
517	G03797	Homo sapiens	Human secreted protein, SEQ ID NO: 7878.	92	40
518	AF151083	Homo sapiens	HSPC249	444	98
519	S80864	Homo sapiens	cytochrome c-like polypeptide	318	50
520	X92485	Plasmodium	pval	170	61
		vivax			ļ

SEQ ID NO:	Accession No.	Species	Description	Smith- Waterman Score	% Identity
521	G03790	Homo sapiens	Human secreted protein, SEQ ID NO: 7871.	159	59
522	AF121857	Homo sapiens	sorting nexin 7	259	40
523	G02654	Homo sapiens	Human secreted protein, SEQ ID NO: 6735.	82	37
524	W88627	Homo sapiens	Secreted protein encoded by gene 94 clone HPMBQ32.	253	73
525	AF119851	Homo sapiens	PRO1722	162	57
526	Y27761	Homo sapiens	Human secreted protein encoded by gene No. 47.	154	57
527	G02707	Homo sapiens	Human secreted protein, SEQ ID NO: 6788.	70	45
528	U47924	Homo sapiens	C8	1112	86
529	G04063	Homo sapiens	Human secreted protein, SEQ ID NO: 8144.	84	45
530	G03203	Homo sapiens	Human secreted protein, SEQ ID NO: 7284.	111	60
531	G04067	Homo sapiens	Human secreted protein, SEQ ID NO: 8148.	92	65
532	G03267	Homo sapiens	Human secreted protein, SEQ ID NO: 7348.	75	29
533	G03203	Homo sapiens	Human secreted protein, SEQ ID NO: 7284.	182	48
534	AF068286	Homo sapiens	HDCMD38P	861	100
535	U07707	Homo sapiens	epidermal growth factor receptor substrate	228	60
536	G01955	Homo sapiens	Human secreted protein, SEQ ID NO: 6036.	484	75
537	AF219232	Gallus gallus	qin-induced kinase	206	53
538	AF135022	Homo sapiens	mediator	128	100
539	G03267	Homo sapiens	Human secreted protein, SEQ ID NO: 7348.	141	59
540	AF016430	Caenorhabditi s elegans	contains similarity to a BR-C/TTK domain	853	39
541	AC003093	Homo sapiens	OXYSTEROL-BINDING PROTEIN; 45% similarity to P22059 (PID:g129308)	408	66
542	M29487	Homo sapiens	integrin alpha subunit precursor	517	81
543	AF102530	Mus musculus	olfactory receptor F3	327	73
544	Y73431	Homo sapiens	Human secreted protein clone yb186_1 protein sequence SEQ ID NO:84.	386	100
545	AE004833	Pseudomonas aeruginosa	probable TonB-dependent receptor	279	42
546	G03793	Homo sapiens	Human secreted protein, SEQ ID NO: 7874.	264	53
547	Y69192	Homo sapiens	A human monocyte-macrophage apolipoprotein B receptor protein.	1772	67
548	Y91493	Homo sapiens	Human secreted protein sequence encoded by gene 43 SEQ ID NO:166.	176	100
549	G01571	Homo sapiens	Human secreted protein, SEQ ID NO: 5652.	777	99
550	AF044588	Homo sapiens	protein regulating cytokinesis 1; PRC1	1953	88
551	Y29332	Homo sapiens	Human secreted protein clone pe584_2 protein sequence.	1224	94
552	X98330	Homo sapiens	ryanodine receptor 2	24621	99
553 554	Y42782	Homo sapiens	Human UC Band #331 protein.	684	95
	AB025258	Mus musculus	granuphilin-a	501	41
555 556	AJ010346	Homo sapiens	RING-H2	1468	100
556 557	W92388 AF119851	Homo sapiens	Human TR-interacting protein S239a. PRO1722	538	92
558	AF117756	Homo sapiens Homo sapiens	thyroid hormone receptor-associated protein	175 183	59 32
559	G02872	Homo sapiens	complex component TRAP150  Human secreted protein, SEQ ID NO: 6953.	319	68
560	D86214	Mus musculus	Ca2+ dependent activator protein for secretion	1010	93
561	AF187325	Canis familiaris	melanoma antigen	287	55
562	AJ001981	Homo sapiens	OXA1L	2512	99
563	Z17238	Rattus norvegicus	glutamate receptor subtype delta-1	338	66
564	W30638	Homo sapiens	Partial human 7-transmembrane receptor HAPO167 protein.	371	100
565	AC005620	Homo sapiens	R33590_1	467	97
566	Y99358	Homo sapiens	Human PRO1772 (UNQ834) amino acid sequence SEQ ID NO:63.	1138	78
567	AL031177	Homo sapiens	dJ889M15,3 (novel protein)	1002	58
568	AF151043	Homo sapiens	HSPC209	798	100

SEQ	Accession	Species	Description	Smith-	%
ID T	No.	F	,	Waterman	Identity
NO:		1		Score	
569	AF097518	Homo sapiens	liver-specific transporter	231	100
570	AB035698	Homo sapiens	Misshapen/NIK-related kinase MINK-1	1532	100
571	Y07096	Homo sapiens	Colon cancer associated antigen precursor sequence.	1064	100
572	AL031177	Homo sapiens	dJ889M15.3 (novel protein)	735	55
573	Y66639	Homo sapiens	Membrane-bound protein PRO290.	254	45
574	AB037108	Homo sapiens	seven transmembrane domain orphan receptor	1883	99
575	D43949	Homo sapiens	This gene is novel.	836	100
576	Y48596	Homo sapiens	Human breast tumour-associated protein 57.	108	50
577	G00352	Homo sapiens	Human secreted protein, SEQ ID NO: 4433.	141	75
578	R95913	Homo sapiens	Neural thread protein.	140	65
579	AK025116	Homo sapiens	unnamed protein product	201	70
580	Y86473	Homo sapiens	Human gene 52-encoded protein fragment, SEQ ID NO:388.	77	70
581	AF196779	Homo sapiens	JM10 protein	450	100
582	AF188706	Homo sapiens	g20 protein	330	98
583	AB030234	Canis familiaris	D4 dopamine receptor	64	56
584	G02621	Homo sapiens	Human secreted protein, SEQ ID NO: 6702.	345	90
585	AL096828	Homo sapiens	dJ963E22.1 (Novel protein similar to NY-REN-2 Antigen)	268	85
586	Y30819	Homo sapiens	Human secreted protein encoded from gene 9.	235	35
587	G00357	Homo sapiens	Human secreted protein, SEQ ID NO: 4438.	132	56
588	G02872	Homo sapiens	Human secreted protein, SEQ ID NO: 6953.	182	79
589	AF235017	Mus musculus	2P1 protein	764	80
590	W88627	Homo sapiens	Secreted protein encoded by gene 94 clone HPMBQ32.	329	81
591	Y30709	Homo sapiens	Amino acid sequence of a human secreted protein.	110	43
592	Y53875	Homo sapiens	A human seven transmembrane signal transducer polypeptide.	1369	92
593	Y53051	Homo sapiens	Human secreted protein clone dd119_4 protein sequence SEQ ID NO:108.	1112	97
594	Y27658	Homo sapiens	Human secreted protein encoded by gene No. 92.	763	79
595	G03798	Homo sapiens	Human secreted protein, SEQ ID NO: 7879.	156	58
596	AF151110	Mus musculus	COP1 protein	2215	95
597	G03786	Homo sapiens	Human secreted protein, SEQ ID NO: 7867.	157	65
598	AF192499	Mus musculus	putative secreted protein ZSIG37	143	40
599	AF119855	Homo sapiens	PRO1847	236	76
600.	G02872	Homo sapiens	Human secreted protein, SEQ ID NO: 6953.	212	73
601	Y00295	Homo sapiens	Human secreted protein encoded by gene 38.	567	88
602	AF184971	Homo sapiens	class II cytokine receptor ZCYTOR7	2015	74
603	AF061936	Homo sapiens	diacylglycerol kinase iota	773	96
604	AL096828	Homo sapiens	dJ963E22.1 (Novel protein similar to NY-REN-2 Antigen)	1333	93
605	AB033106	Homo sapiens	KIAA1280 protein	3915	100
606	X75756	Homo sapiens	protein kinase C mu	3916	99
607	D86983	Homo sapiens	similar to D.melanogaster peroxidasin(U11052)	5758	99
608	W69341	Homo sapiens	Secreted protein of clone CG279_1.	1377	99
609	W88627	Homo sapiens	Secreted protein encoded by gene 94 clone HPMBQ32.	339	82
610	Y27868	Homo sapiens	Human secreted protein encoded by gene No. 107.	116	62
611	AF202636	Homo sapiens	angiopoietin-like protein PP1158	2164	100
612	AF090944	Homo sapiens	PRO0663	218	82
613	Y02693	Homo sapiens	Human secreted protein encoded by gene 44 clone HTDAD22.	195	59
614	M87053	Rattus norvegicus	lens membrane protein	450	84
615	AC004232	Homo sapiens	FPM315	163	37
616	G01984	Homo sapiens	Human secreted protein, SEQ ID NO: 6065.	205	79

SEQ ID NO:	Accession No.	Species	Description	Smith- Waterman Score	% Identity
617	Y91524	Homo sapiens	Human secreted protein sequence encoded by gene 74 SEQ ID NO:197.	821	99
618	AJ245621	Homo sapiens	CTL2 protein	2258	99
619	Y76198	Homo sapiens	Human secreted protein encoded by gene 75.	108	64
620	AF067864	Homo sapiens	transferrin receptor 2 alpha	3922	94
621	D90721	Escherichia coli	Transmembrane protein dppC	573	90
622	W75858	Homo sapiens	Human secretory protein of clone CS752-3.	730	100
623	Y94982	Homo sapiens	Human secreted protein vb12 1, SEQ ID NO:4.	733	100
624	AF034745	Mus musculus	LNXp80	637	83
625	U42580	Paramecium bursaria Chlorella virus 1	Pro-rich, IPPPNMSLPLS (3x)	94	46
626	U79260	Homo sapiens	unknown	194	70
627	R95913	Homo sapiens	Neural thread protein.	99	50
628	G03450	Homo sapiens	Human secreted protein, SEQ ID NO: 7531.	427	100
629	Y36281	Homo sapiens	Human secreted protein encoded by gene 58.	590	100
630	Y02693	Homo sapiens	Human secreted protein encoded by gene 44 clone HTDAD22.	165	76
631	G02139	Homo sapiens	Human secreted protein, SEQ ID NO: 6220.	268	96
632	U16996	Homo sapiens	protein tyrosine posphatase	351	80
633	AF121857	Homo sapiens	sorting nexin 7	2019	100
634	AF283772	Homo sapiens	similar to Homo sapiens ribosomal protein L10 encoded by GenBank Accession Number L25899	340	77
635	Y07090	Homo sapiens	Renal cancer associated antigen precursor sequence.	277	64
636	AB013382	Homo sapiens	DÚSP6	414	76
637	G02872	Homo sapiens	Human secreted protein, SEQ ID NO: 6953.	315	71
638	M95762	Rattus norvegicus	GABA transporter	924	89
639	G03789	Homo sapiens	Human secreted protein, SEQ ID NO: 7870.	219	60
640	Y01400	Homo sapiens	Secreted protein encoded by gene 18 clone HNHFO29.	137	79
641	AC008075	Arabidopsis thaliana	F24J5.4	121	33
642	W74824	Homo sapiens	Human secreted protein encoded by gene 96 clone HAQBK61.	615	62
643	AB015982	Homo sapiens	serine/threonine kinase	485	98
644	Y25806	Homo sapiens	Human secreted protein fragment encoded from gene 23.	162	46
645	AF122904	Homo sapiens	membrane protein DAP10	474	100
646	AF233323	Homo sapiens	Fas-associated phosphatase-1	200	38
647	W48804	Homo sapiens	Homo sapiens clone BK158 1 protein.	1203	99
648	AF257330	Homo sapiens	COBW-like protein	1440	98
549	Y36203	Homo sapiens	Human secreted protein #75.	233	73
650	G02872		Human secreted protein, SEQ ID NO: 6953.		
651	Y32199	Homo sapiens Homo sapiens	Human receptor molecule (REC) encoded by	173 1012	78 100
652	AB032909	Hylobates agilis	Incyte clone 2022379. dopamine receptor D4	122	32
653	AK021848	Homo sapions	unnamed protein product	186	69
654	W73411	Homo sapiens	Human secreted protein encoded by Gene No. 15.	57	37
655	L22455	Rattus	mu opioid receptor	116	34
656	G03112	Homo sapiens	Human secreted protein, SEQ ID NO: 7193.	110	45
557	G02345	Homo sapiens	Human secreted protein, SEQ ID NO: 7193.  Human secreted protein, SEQ ID NO: 6426.	459	97
658	W88627	Homo sapiens	Secreted protein encoded by gene 94 clone HPMBQ32.	291	75
659	G02832	Homo sapiens	Human secreted protein, SEQ ID NO: 6913.	134	65
660	Y91423	Homo sapiens	Human secreted protein sequence encoded by	333	96
	1.2.		gene 11 SEQ ID NO:144.		

SEQ ID NO:	Accession No.	Species	Description	Smith- Waterman Score	% Identity
661	G03789	Homo sapiens	Human secreted protein, SEQ ID NO: 7870.	168	68 .
662	Y53886	Homo sapiens	A suppressor of cytokine signalling protein designated HSCOP-6.	375	43
663	W75771	Homo sapiens	Human GTP binding protein APD08.	629	100
664	AL096770	Homo sapiens	bA150A6.2 (novel 7 transmembrane receptor (rhodopsin family) (olfactory receptor like) protein (hs6M1-21))	480	55
665	AB037734	Homo sapiens	KIAA1313 protein	978	96
666	W82841	Homo sapiens	Human cerebral protein-1.	192	84
667	W82841	Homo sapiens	Human cerebral protein-1.	182	87
668	AB030184	Mus musculus	contains transmembrane (TM) region and ATP binding region	757	68
669	AB032919	Hylobates muelleri	dopamine receptor D4	85	37
670	AF107295	Rattus norvegicus	outer membrane protein	746	81
671	Z33642	Homo sapiens	leukocyte surface protein	394	93
672	W85608	Homo sapiens	Secreted protein clone du410_5.	261	91
673	G03203	Homo sapiens	Human secreted protein, SEQ ID NO: 7284.	106	48
674	AL035587	Homo sapiens	dJ475N16.4 (KIAA0240)	2388	99
675 676	Y59668 G03797	Homo sapiens	Secreted protein 108-005-5-0-C1-FL.	1134 174	53
677	AF026954	Homo sapiens Bos taurus	Human secreted protein, SEQ ID NO: 7878.  pyruvate dehydrogenase phosphatase regulatory	1013	74 95
			subunit precursor; PDPr		
678	L11625	Mus musculus	receptor protein-tyrosine kinase	545	96
679	AL031427	Homo sapiens	dJ167A19.3 (novel protein)	745	100
680	AJ133430	Mus musculus	olfactory receptor	528	77
681	G02532	Homo sapiens	Human secreted protein, SEQ ID NO: 6613.	179	70
682	G03789	Homo sapiens	Human secreted protein, SEQ ID NO: 7870.	336	76
683	Y94943	Homo sapiens	Human secreted protein clone yt14_1 protein sequence SEQ ID NO:92.	118	100
684	U43360	Peromyscus maniculatus	reverse transcriptase	100	37
685	G00885	Homo sapiens	Human secreted protein, SEQ ID NO: 4966.	162	60
686	AK001518	Homo sapiens	unnamed protein product	590	100
687	G01982	Homo sapiens	Human secreted protein, SEQ ID NO: 6063.	718	100
688	Y92241	Homo sapiens	Human cancer associated antigen precursor (MO-REN-46).	2405	99
689	AC024792	Caenorhabditi s elegans	contains similarity to TR:P78316	423	36
690	Y27868	Homo sapiens	Human secreted protein encoded by gene No. 107.	183	81
691	Y56514	Homo sapiens	Human Jurkat cell clone P2-15 AIM10 longest ORF protein sequence.	180	88
692	Y27795	Homo sapiens	Human secreted protein encoded by gene No. 79.	1539	99
693	Y36268	Homo sapiens	Human secreted protein encoded by gene 45.	428	98
694	U12465	Homo sapiens	ribosomal protein L35	308	89
695	Y45272	Homo sapiens	Human secreted protein encoded from gene 16.	1517	99
696	AF191838	Homo sapiens	TANK binding kinase TBK1	1242	98
697	Y02693	Homo sapiens	Human secreted protein encoded by gene 44 clone HTDAD22.	275	75
698	Y87280	Homo sapiens	Human signal peptide containing protein HSPP57 SEQ ID NO:57.	576	90
699	Y97999	Homo sapiens	Human SCAD family molecule HSFM-1, SEQ ID NO:1.	729	99
700	AJ006701	Homo sapiens	putative serine/threonine protein kinase	610	79
701	AF209198	Homo sapiens	zinc finger protein 277	2357	100
702	AJ298841	Mus musculus	torsinA protein	709	45
703	AK021729	Homo sapiens	unnamed protein product	622	98
704	Z46787	Caenorhabditi s elegans	similar to Glutaredoxin, Zinc finger, C3HC4 type (RING finger)	920	51
705	G02882	Homo sapiens	Human secreted protein, SEQ ID NO: 6963.	589	98

SEQ	Accession	Species	Description	Smith-	%
ID	No.	,		Waterman	Identity
NO:	-	<u> </u>		Score	<u> </u>
706	G02501	Homo sapiens	Human secreted protein, SEQ ID NO: 6582.	125	58
707	R95326	Homo sapiens	Tumor necrosis factor receptor 1 death domain	121	95
700	002000	7.	ligand (clone 2DD).	105	ļ
708	G03002	Homo sapiens	Human secreted protein, SEQ ID NO: 7083.	125	39
709 710	Y96202 M63577	Homo sapiens	IkappaB kinase (IKK) binding protein, Y2H56.	516	98
/10	M033//	Saccharomyc es cerevisiae	SFP1	131	59
711	AB026291	Rattus	acetoacetyl-CoA synthetase	467	85
/ 1 1	AD020271	norvegicus	accidactlyi-coa symmetase	407	6.5
712	D21211	Homo sapiens	protein tyrosine phosphatase (PTP-BAS, type 3)	368	44
713	AF044033	Marmota	olfactory receptor	615	83
	12011000	marmota	onactory receptor	015	03
714	G03561	Homo sapiens	Human secreted protein, SEQ ID NO: 7642.	251	100
715	AB033062	Homo sapiens	KIAA1236 protein	1380	100
716	G00577	Homo sapiens	Human secreted protein, SEQ ID NO: 4658.	80	73
717	Y96864	Homo sapiens	SEQ. ID. 37 from WO0034474.	835	99
718	AJ243396	Homo sapiens	voltage-gated sodium channel beta-3 subunit	234	100
719	U47334	Homo sapiens	similar to chicken gamma aminobutyric acid	578	99
			receptor beta4 subunit	I	1
720	AB020598	Homo sapiens	peptide transporter 3	1096	100
721	Y53886	Homo sapiens	A suppressor of cytokine signalling protein	570	74
			designated HSCOP-6.		1_
722	J05046	Homo sapiens	insulin receptor-related receptor	6787	100
723	AF001958	Ambystoma	electrogenic Na+ bicarbonate cotransporter;	111	41
		tigrinum	NBC	L	
724	AF127084	Mus	semaphorin cytoplasmic domain-associated	5253	94
		musculus	protein 3A		
725	X54673	Homo sapiens	GABA transporter	3114	99
726	AF016191	Rattus	potassium channel	370	100
705	1.0000000	norvegicus			
727	AB029559	Rattus	BATI	139	35
700	V20502	norvegicus	HOEH2 H	2106	0.5
728 729	Y28503 AJ011415	Homo sapiens	HGFH3 Human Growth Factor Homologue 3.	2186	97
730	Z93096	Homo sapiens Homo sapiens	plexin-B1/SEP receptor bK390B3.1 (manic fringe (Drosophila)	729 142	56 68
130	2,93090	nomo sapiens	homolog)	142	00
731	Z10062	Homo sapiens	cDNA encoding a human vanilloid receptor	675	99
751	210002	Home sapiens	homologue Vanilrep1.	073	) 33
732	AF161382	Homo sapiens	HSPC264	492	94
733	AB029033	Homo sapiens	KIAA1110 protein	3826	99
734	AE000493	Escherichia	putative transport protein	592	97
		coli	parative manopoli protoni	1 332	- '
735	AL033379	Homo sapiens	dJ417O22.2 (novel 7 transmembrane receptor	2173	99
			(rhodopsin family) protein similar to high-		1
	<u> </u>		affinity lysophosphatidic acid receptor homolog)		
736	AF132599	Homo sapiens	RANTES factor of late activated T lymphocytes-	245	56
	<u> </u>		1		
737	X55019	Homo sapiens	acetylcholine receptor delta subunit	883	99.
738	X91906	Homo sapiens	voltage-gated chloride ion channel	1978	100
739	AB026116	Homo sapiens	organic anion transporter 4	1444	98
740	D00570	Mus	open reading frame (196 AA)	83	24
	<u> </u>	musculus			
741	W03626	Homo sapiens	Human thyrotropin GPR N-terminal sequence.	118	40
742	U66059	Homo sapiens	V_segment translation product	614	100
743	AF119815	Homo sapiens	G-protein-coupled receptor	2751	99
744	X16663	Homo sapiens	haematopoietic lineage cell protein (AA 1-486)	148	93
745	W67838	Homo sapiens	Human secreted protein encoded by gene 32	448	95
746	WEGOCO	1,77	clone HLTCJ63.	0414	100
746	W57260	Homo sapiens	Human semaphorin Y.	2414	100
747	W21578	Homo sapiens	Alzheimer's disease protein encoded by DNA	968	65
748	V04025	Homo comics -	from plasmid pGCS2232.	622	100
740	Y94935	Homo sapiens	Human secreted protein clone yd218_1 protein sequence SEQ ID NO:76.	622	100
		_1			
749	AL022238	Homo sapiens	dJ1042K10.5 (novel protein)	314	85

AB025258   Mus musculus   gramphilin-a   773   41	SEQ ID NO:	Accession No.	Species	Description	Smith- Waterman Score	% Identity
		AB025258		granuphilin-a		41
Y48586	752	Y52386		Human transmembrane protein HP02000.	900	99
Main						
M85183				nutative G protein coupled recentor 97		
Procedure   Proc			norvegicus			
			•	receptor 6		
R04932	757					
39 responsible for binding the target.   99	758	Z22535	Homo sapiens	ALK-3	439	98
Homo sapiens	759	R04932	Homo sapiens	Interferon-gamma receptor segment from clone 39 responsible for binding the target.	564	97
AB020676   Homo sapiens   KIAA0869 protein   4433   99	760	W74902	Homo sapiens	Human secreted protein encoded by gene 175	1217	99
AB020676   Homo sapiens   KIAA0869 protein   4433   99	761	G03706	Homo sapiens	Human secreted protein, SEO ID NO: 7787.	223	88
763         AK026992         Homo sapiens         unnamed protein product         2285         99           764         AF173358         Homo sapiens         glucocorticoid receptor AF-1 coactivator-1         573         100           765         AF268066         Mus musculus         netrin 4         2019         89           766         Y48585         Homo sapiens         Human breast tumour-associated protein 46.         1169         89           767         AF230378         Mus musculus         dorant receptor S18         268         62           768         AF121975         Mus musculus         dorant receptor S18         268         62           769         AB008515         Homo sapiens         Raffus         putative integral membrane transport protein         458         50           770         Y09945         Raffus         putative integral membrane transport protein         458         50           771         AF226731         Homo sapiens         HOW         Putative integral membrane transport protein         458         50           771         AF226731         Homo sapiens         HOW         Putative integral membrane transport protein         458         50           777         AF125101         Homo sapiens         Homo sa						
AF173358						
AF268066   Mus   musculus   musculus   musculus   Mus   musculus   musculus						I
Mus						
AF230378			musculus			
Musculus		Homo sapiens		1		
768         AF121975         Mus musculus musculus musculus         odorant receptor S18 musculus         268         62           769         AB008515         Homo sapiens         RanBPM         611         57           770         Y09945         Rattus norvegicus         putative integral membrane transport protein         458         50           771         AF226731         Homo sapiens         AD026         688         99           772         Y27132         Homo sapiens         NOV/plexin-A1 protein         1821         98           773         X87832         Homo sapiens         NOV/plexin-A1 protein         1821         98           774         AB025258         Mus musculus         granuphilin-a         500         41           775         AF125101         Homo sapiens         HSPC040 protein         232         93           776         G02815         Homo sapiens         Human secreted protein, SEQ ID NO: 6594.         191         68           777         G02493         Homo sapiens         Sequence of pre-human atrial natriuretic peptide.         213         45           778         R03301         Homo sapiens         Sequence of pre-human atrial natriuretic peptide.         213         45           780	767	AF230378		interleukin-1 delta	309	45
Rabous   R	768	AF121975		odorant receptor S18	268	62
Possible    769	AB008515		RanBPM	611	57	
AF226731			Rattus			
Human glioblastoma-derived polypeptide (clone OA004FG).   1384   100 OA004FG).   1384   100 OA004FG).   1384   100 OA004FG).   13821	771	A F226721		AD026	600	100
773         X87832         Homo sapiens         NOV/plexin-A1 protein         1821         98           774         AB025258         Mus musculus         granuphilin-a         500         41           775         AF125101         Homo sapiens         HSPC040 protein         232         93           776         G02815         Homo sapiens         Human secreted protein, SEQ ID NO: 6896.         314         95           777         G02493         Homo sapiens         Human secreted protein, SEQ ID NO: 6896.         314         95           778         R03301         Homo sapiens         Sequence of pre-human atrial natriuretic peptide.         213         45           779         AL357374         Homo sapiens         Sequence of pre-human atrial natriuretic peptide.         213         45           780         AF100346         Homo sapiens         neuronal voltage gated calcium channel gamma-         1434         89           781         Y19566         Homo sapiens         Human secreted protein encoded by gene 10.         1098         93           782         Y36233         Homo sapiens         Human low density lipoprotein binding protein         2693         99           783         AF084464         Rattus         GTP-binding protein REM2         141		(		Human glioblastoma-derived polypeptide (clone	1	1
AB025258	773	X87832	Homo saniens		1821	98
775         AF125101         Homo sapiens         HSPC040 protein         232         93           776         G02815         Homo sapiens         Human secreted protein, SEQ ID NO: 6896.         314         95           777         G02493         Homo sapiens         Human secreted protein, SEQ ID NO: 6574.         191         68           778         R03301         Homo sapiens         Sequence of pre-human atrial natriuretic peptide.         213         45           779         AL357374         Homo sapiens         bA353C18.2 (novel protein)         232         100           780         AF100346         Homo sapiens         neuronal voltage gated calcium channel gamma-3 subunit         1434         89           781         Y19566         Homo sapiens         Amino acid sequence of a human secreted         103         52           782         Y36233         Homo sapiens         Human secreted protein encoded by gene 10.         1098         93           783         AF084464         Rattus         GTP-binding protein REM2         141         30           784         W49042         Homo sapiens         Human low density lipoprotein binding protein         2693         99           785         AF238381         Homo sapiens         Human apoptosis related protein. <td></td> <td></td> <td>Mus</td> <td>granuphilin-a</td> <td></td> <td></td>			Mus	granuphilin-a		
776         G02815         Homo sapiens         Human secreted protein, SEQ ID NO: 6896.         314         95           777         G02493         Homo sapiens         Human secreted protein, SEQ ID NO: 6574.         191         68           778         R03301         Homo sapiens         Sequence of pre-human atrial natriuretic peptide.         213         45           779         AL357374         Homo sapiens         bA353C18.2 (novel protein)         232         100           780         AF100346         Homo sapiens         neuronal voltage gated calcium channel gamma-3 subunit         1434         89           781         Y19566         Homo sapiens         Amino acid sequence of a human secreted protein accoded by gene 10.         1098         93           782         Y36233         Homo sapiens         Human secreted protein encoded by gene 10.         1098         93           783         AF084464         Rattus         GTP-binding protein REM2         141         30           784         W49042         Homo sapiens         Human low density lipoprotein binding protein         2693         99           785         AF238381         Homo sapiens         Human apoptosis related protein.         547         100           786         Y91870         Homo sapiens	THE	AE125101		LICECOMO	1 222	102
Homo sapiens   Human secreted protein, SEQ ID NO: 6574.   191   68   68   68   68   68   68   69   69						
778         R03301         Homo sapiens         Sequence of pre-human atrial natriuretic peptide.         213         45           779         AL357374         Homo sapiens         bA353C18.2 (novel protein)         232         100           780         AF100346         Homo sapiens         neuronal voltage gated calcium channel gamma-3 subunit         1434         89           781         Y19566         Homo sapiens         Amino acid sequence of a human secreted protein.         103         52           782         Y36233         Homo sapiens         Human secreted protein encoded by gene 10.         1098         93           783         AF084464         Rattus norvegicus         GTP-binding protein REM2         141         30           784         W49042         Homo sapiens         Human low density lipoprotein binding protein         2693         99           785         AF238381         Homo sapiens         Human apoptosis related protein.         547         100           786         Y91870         Homo sapiens         Human membrane transport protein, MTRP-7.         1062         94           787         Y71062         Homo sapiens         Human membrane transport protein, MTRP-7.         1062         94           789         AL049569         Homo sapiens						
AL357374   Homo sapiens   bA353C18.2 (novel protein)   232   100			Homo sapiens	Human secreted protein, SEQ ID NO: 6574.		
Name			Homo sapiens	Sequence of pre-human atrial natriuretic peptide.		
3 subunit   103   52	779	AL357374		bA353C18.2 (novel protein)	232	100
Protein.   Protein.	780	AF100346	Homo sapiens	neuronal voltage gated calcium channel gamma-	1434	89
782         Y36233         Homo sapiens         Human secreted protein encoded by gene 10.         1098         93           783         AF084464         Rattus norvegicus         GTP-binding protein REM2         141         30           784         W49042         Homo sapiens         Human low density lipoprotein binding protein         2693         99           785         AF238381         Homo sapiens         PTOV1         1904         91           786         Y91870         Homo sapiens         Human apoptosis related protein.         547         100           787         Y71062         Homo sapiens         Human membrane transport protein, MTRP-7.         1062         94           788         AF117754         Homo sapiens         thyroid hormone receptor-associated protein         8684         98           789         AL049569         Homo sapiens         dJ37C10.3 (novel ATPase)         2848         96           790         AF151848         Homo sapiens         CGI-90 protein         745         96           791         Y08639         Homo sapiens         Human PRO381 protein sequence.         644         99           793         AF121228         Homo sapiens         thyroid hormone receptor-associated protein         1037         100	781	Y19566	Homo sapiens	I	103	52
AF084464   Rattus   norvegicus   Rem2   Rem2   Rem2   Rem3   Re	782	Y36233	Homo saniens		1098	93
784         W49042         Homo sapiens         Human low density lipoprotein binding protein LBP-3.         2693         99           785         AF238381         Homo sapiens         PTOV1         1904         91           786         Y91870         Homo sapiens         Human apoptosis related protein.         547         100           787         Y71062         Homo sapiens         Human membrane transport protein, MTRP-7.         1062         94           788         AF117754         Homo sapiens         thyroid hormone receptor-associated protein         8684         98           789         AL049569         Homo sapiens         dJ37C10.3 (novel ATPase)         2848         96           790         AF151848         Homo sapiens         CGI-90 protein         745         96           791         Y08639         Homo sapiens         nuclear orphan receptor ROR-beta         1421         95           792         Y41706         Homo sapiens         Human PRO381 protein sequence.         644         99           793         AF121228         Homo sapiens         thyroid hormone receptor-associated protein         1037         100           794         G04072         Homo sapiens         Human secreted protein, SEQ ID NO: 8153.         124         62 </td <td></td> <td></td> <td>Rattus</td> <td></td> <td></td> <td></td>			Rattus			
785         AF238381         Homo sapiens         PTOV1         1904         91           786         Y91870         Homo sapiens         Human apoptosis related protein.         547         100           787         Y71062         Homo sapiens         Human membrane transport protein, MTRP-7.         1062         94           788         AF117754         Homo sapiens         thyroid hormone receptor-associated protein         8684         98           789         AL049569         Homo sapiens         dJ37C10.3 (novel ATPase)         2848         96           790         AF151848         Homo sapiens         CGI-90 protein         745         96           791         Y08639         Homo sapiens         nuclear orphan receptor ROR-beta         1421         95           792         Y41706         Homo sapiens         Human PRO381 protein sequence.         644         99           793         AF121228         Homo sapiens         thyroid hormone receptor-associated protein         1037         100           complex component TRAP95         complex component TRAP95         124         62           795         Y69384         Homo sapiens         Amino acid sequence of a 14274 receptor         119         100	784	W49042		, , , , , ,	2693	99
786         Y91870         Homo sapiens         Human apoptosis related protein.         547         100           787         Y71062         Homo sapiens         Human membrane transport protein, MTRP-7.         1062         94           788         AF117754         Homo sapiens         thyroid hormone receptor-associated protein         8684         98           789         AL049569         Homo sapiens         dJ37C10.3 (novel ATPase)         2848         96           790         AF151848         Homo sapiens         CGI-90 protein         745         96           791         Y08639         Homo sapiens         nuclear orphan receptor ROR-beta         1421         95           792         Y41706         Homo sapiens         Human PRO381 protein sequence.         644         99           793         AF121228         Homo sapiens         thyroid hormone receptor-associated protein         1037         100           complex component TRAP95         complex component TRAP95         124         62           795         Y69384         Homo sapiens         Amino acid sequence of a 14274 receptor         119         100	785	AF238381	Homo seniens		1904	91
787         Y71062         Homo sapiens         Human membrane transport protein, MTRP-7.         1062         94           788         AF117754         Homo sapiens         thyroid hormone receptor-associated protein         8684         98           789         AL049569         Homo sapiens         dJ37C10.3 (novel ATPase)         2848         96           790         AF151848         Homo sapiens         CGI-90 protein         745         96           791         Y08639         Homo sapiens         nuclear orphan receptor ROR-beta         1421         95           792         Y41706         Homo sapiens         Human PRO381 protein sequence.         644         99           793         AF121228         Homo sapiens         thyroid hormone receptor-associated protein complex component TRAP95         1037         100           794         G04072         Homo sapiens         Human secreted protein, SEQ ID NO: 8153.         124         62           795         Y69384         Homo sapiens         Amino acid sequence of a 14274 receptor         119         100						
788         AF117754         Homo sapiens         thyroid hormone receptor-associated protein complex component TRAP240         8684         98           789         AL049569         Homo sapiens         dJ37C10.3 (novel ATPase)         2848         96           790         AF151848         Homo sapiens         CGI-90 protein         745         96           791         Y08639         Homo sapiens         nuclear orphan receptor ROR-beta         1421         95           792         Y41706         Homo sapiens         Human PRO381 protein sequence.         644         99           793         AF121228         Homo sapiens         thyroid hormone receptor-associated protein complex component TRAP95         1037         100           794         G04072         Homo sapiens         Human secreted protein, SEQ ID NO: 8153.         124         62           795         Y69384         Homo sapiens         Amino acid sequence of a 14274 receptor         119         100						
Complex component TRAP240						
790         AF151848         Homo sapiens         CGI-90 protein         745         96           791         Y08639         Homo sapiens         nuclear orphan receptor ROR-beta         1421         95           792         Y41706         Homo sapiens         Human PRO381 protein sequence.         644         99           793         AF121228         Homo sapiens         thyroid hormone receptor-associated protein complex component TRAP95         1037         100           794         G04072         Homo sapiens         Human secreted protein, SEQ ID NO: 8153.         124         62           795         Y69384         Homo sapiens         Amino acid sequence of a 14274 receptor         119         100				complex component TRAP240		
790         AF151848         Homo sapiens         CGI-90 protein         745         96           791         Y08639         Homo sapiens         nuclear orphan receptor ROR-beta         1421         95           792         Y41706         Homo sapiens         Human PRO381 protein sequence.         644         99           793         AF121228         Homo sapiens         thyroid hormone receptor-associated protein complex component TRAP95         1037         100           794         G04072         Homo sapiens         Human secreted protein, SEQ ID NO: 8153.         124         62           795         Y69384         Homo sapiens         Amino acid sequence of a 14274 receptor         119         100			Homo sapiens		1	3
791         Y08639         Homo sapiens         nuclear orphan receptor ROR-beta         1421         95           792         Y41706         Homo sapiens         Human PRO381 protein sequence.         644         99           793         AF121228         Homo sapiens         thyroid hormone receptor-associated protein complex component TRAP95         1037         100           794         G04072         Homo sapiens         Human secreted protein, SEQ ID NO: 8153.         124         62           795         Y69384         Homo sapiens         Amino acid sequence of a 14274 receptor         119         100	790	AF151848			745	96
792         Y41706         Homo sapiens         Human PRO381 protein sequence.         644         99           793         AF121228         Homo sapiens         thyroid hormone receptor-associated protein complex component TRAP95         1037         100           794         G04072         Homo sapiens         Human secreted protein, SEQ ID NO: 8153.         124         62           795         Y69384         Homo sapiens         Amino acid sequence of a 14274 receptor         119         100						
793         AF121228         Homo sapiens         thyroid hormone receptor-associated protein complex component TRAP95         1037         100           794         G04072         Homo sapiens         Human secreted protein, SEQ ID NO: 8153.         124         62           795         Y69384         Homo sapiens         Amino acid sequence of a 14274 receptor         119         100						
794         G04072         Homo sapiens         Human secreted protein, SEQ ID NO: 8153.         124         62           795         Y69384         Homo sapiens         Amino acid sequence of a 14274 receptor         119         100				thyroid hormone receptor-associated protein		
795 Y69384 Homo sapiens Amino acid sequence of a 14274 receptor 119 100	704	G04072	Homo coniona		124	62
protein.				Amino acid sequence of a 14274 receptor		
796 W40215 Homo sapiens Human macrophage antigen. 1358 99					L	

SEQ	Accession	Species	Description	Smith-	%
ID	No.			Waterman	Identity
NO:	1			Score	
797	AF258340	Homo sapiens	hepatocellular carcinoma-associated antigen 112	1151	99
798	AF159615	Homo sapiens	FGF receptor activating protein 1	461	98
799	Y59863	Homo sapiens	Human normal uterus tissue derived protein 26.	797	99
800	W70459	Homo sapiens	Human T1-receptor ligand III splice variant 2,	572	92
801	L00073	Homo sapiens	renin	1913	93
802	P92219	Homo sapiens	CR1 protein.	11963	97
000	771.50.55	(human)	1272	-	
803	X15357	Homo sapiens	ANP-A receptor preprotein (AA -32 to 1029)	5199	98
804	W64473	Homo sapiens	Human secreted protein from clone EC172_1.	4018	95
805	AJ243874	Homo sapiens	oligophrenin-4	2067	100
806	G01731	Homo sapiens	Human secreted protein, SEQ ID NO: 5812.	284	100
807	Z24680	Homo sapiens	garp	1562	83
808	AF171669	Homo sapiens	glycoprotein-associated amino acid transporter LAT2	1364	90
809	W70321	Homo sapiens	Secreted protein CC198 1.	1154	96
810	W74843	Homo sapiens	Human secreted protein encoded by gene 115	855	99
		1	clone HOVBA03.		
811	AF108831	Homo sapiens	K:Cl cotransporter 3	4561	100
812	AF092135	Homo sapiens	PTD014	862	100
813	AF283772	Homo sapiens	similar to Homo sapiens ribosomal protein L10	784	100
		,	encoded by GenBank Accession Number	}	
			L25899		
814	G01563	Homo sapiens	Human secreted protein, SEQ ID NO: 5644.	330	100
815	AF051151	Homo sapiens	Toll/interleukin-1 receptor-like protein 3	3850	99
816	W95630	Homo sapiens	Homo sapiens secreted protein gene clone	358	100
	1		gn114_1.	ľ	
817	G01082	Homo sapiens	Human secreted protein, SEQ ID NO: 5163.	549	100
818	AF151800	Homo sapiens	CGI-41 protein	1106	95
819	L00352	Homo sapiens	low density lipoprotein receptor	3980	100
820	X04434	Homo sapiens	IGF-I receptor	5832	99
821	G03844	Homo sapiens	Human secreted protein, SEQ ID NO: 7925.	572	100
822	AF212220	Homo sapiens	TERA	396	48
823	Y50125	Homo sapiens	Human glycophosphatidylinositol-anchored	4897	99
824	AF156778	Homo sapiens	protein GPI-122.	2675	98
825	AF096322	Homo sapiens	ASB-3 protein neuronal voltage-gated calcium channel gamma-	1105	100
023	AF090322	nomo sapiens	2 subunit	1103	100
826	Y07972	Homo sapiens	Human secreted protein fragment #2 encoded	1540	100
		•	from gene 28.	1	1
827	AB032013	Homo sapiens	potassium channel Kv8.1	2435	95
828	Y13620	Homo sapiens	BCL9	5284	96
829	Y91474	Homo sapiens	Human secreted protein sequence encoded by	541	98
			gene 24 SEQ ID NO:147.		<u> </u>
830	X54232	Homo sapiens	glypican	1625	87
831	X14830	Homo sapiens	acetylcholine receptor beta-subunit preprotein	2540	100
832	Y71262 .	Homo sapiens	Human chondromodulin-like protein, Zchm1.	1002	100
833	G03873	Homo sapiens	Human secreted protein, SEQ ID NO: 7954.	638	96
834	AC003030	Homo sapiens	R29828_1	1389	93
835	Y38422	Homo sapiens	Human secreted protein.	964	87
836	U41557	Caenorhabditi	glycine-rich	85	36
837	AL121889	s elegans Homo sapiens	dJ1076E17.1 (KIAA0823 protein (continues in	998	75
			AL023803))	l	
838	AJ011415	Homo sapiens	plexin-B1/SEP receptor	1580	60
839	W80398	Homo sapiens	A secreted protein encoded by clone cw1543_3.	1105	67
840	G00862	Homo sapiens	Human secreted protein, SEQ ID NO: 4943.	255	92
841	G02650	Homo sapiens	Human secreted protein, SEQ ID NO: 6731.	644	97
842	AF036717	Homo sapiens	FGFR signalling adaptor SNT-1	2629	99
843	¥73446	Homo sapiens	Human secreted protein clone yc27_1 protein	1089	100
844	G02872	Homo sapiens	sequence SEQ ID NO:114.  Human secreted protein, SEQ ID NO: 6953.	357	69
844	AF151810	Homo sapiens Homo sapiens		1443	88
845 846	X83378	Homo sapiens	CGI-52 protein putative chloride channel	1620	99
040	1000010	rionio sapiens	similar to general transcription factor 2I; similar	655	フプ

SEQ	Accession	Species	Description	Smith-	1%
ID	No.			Waterman	Identity
NO:				Score	
			to AF038969 (PID:g2827207)		
848	X99886	Homo sapiens	monocyte chemotactic protein-2	160	76
849	AC005587	Homo sapiens	similar to mouse olfactory receptor 13; similar to P34984 (PID:g464305)	963	98
850	AB038237	Homo sapiens	G protein-coupled receptor C5L2	1767	100
851	AF124490	Homo sapiens	ARF GTPase-activating protein GIT1	3415	98
852	Y86217	Homo sapiens	Human secreted protein HWHGU54, SEQ ID NO:132.	1189	99
853	AF224741	Homo sapiens	chloride channel protein 7	3748	99
854	X17094	Homo sapiens	furin (AA 1-794)	3550	99
855	W78245	Homo sapiens	Fragment of human secreted protein encoded by gene 19.	1245	99
856	R97569	Homo sapiens	Interleukin-2 receptor associated protein p43.	1926	100
857	Y41765	Homo sapiens	Human PRO1083 protein sequence.	3211	99
858	AF057306	Homo sapiens	transmembrane proteolipid	481	84
859	AK025116	Homo sapiens	unnamed protein product	374	69
860	Y41312	Homo sapiens	Human secreted protein encoded by gene 5 clone HLDRM43.	824	100
862	Y25776	Homo sapiens	Human secreted protein encoded from gene 66.	895	99
863	Y74188	Homo sapiens	Human prostate tumor EST fragment derived protein #375.	96	30
864	AF167473	Homo sapiens	heme-binding protein	870	99
865	G02532	Homo sapiens	Human secreted protein, SEQ ID NO: 6613.	211	67
866	X54870	Homo sapiens	Type II integral membrane protein	1201	100
867	G00700	Homo sapiens	Human secreted protein, SEQ ID NO: 4781.	640	99
868	Y07894	Homo sapiens	Human secreted protein fragment encoded from gene 43.	388	88
869	J00123	Homo sapiens	preproenkephalin (	1349	95
870	Y91632	Homo sapiens	Human secreted protein sequence encoded by gene 25 SEQ ID NO:305.	1048	98
871	L04311	Homo sapiens	GABA-alpha receptor beta-3 subunit	237	93
872	Y29988	Homo sapiens	Human cytokine family member EF-7 protein.	960	94
873	AF161382	Homo sapiens	HSPC264	1124	99
874	G03412	Homo sapiens	Human secreted protein, SEQ ID NO: 7493.	464	100
875	Y27572	Homo sapiens	Human secreted protein encoded by gene No. 6.	573	96
876	M15530	Homo sapiens	B-cell growth factor	171	56
877 878	W63681 L27867	'Homo sapiens Rattus	Human secreted protein 1. neurexophilin	1652 1448	99
879	Y10835	norvegicus Homo sapiens	Amino acid sequence of a human secreted	321	100
-	,		protein.		
880	W88991	Homo sapiens	Polypeptide fragment encoded by gene 144.	936	100
881	AF118670	Homo sapiens	orphan G protein-coupled receptor	1971	100
882	AF208865	Homo sapiens	EDRF	528	100
883 884	Y18462 Y94950	Homo sapiens Homo sapiens	cathepsin L Human secreted protein clone dh1073_12 protein sequence SEQ ID NO:106.	209 348	72 100
885	AF070661	Homo sapiens	HSPC005	404	100
886	Y04315	Homo sapiens	Human secreted protein encoded by gene 23.	385	100
887	X92744	Homo sapiens	hBD-1	375	100
888	Y22496	Homo sapiens	Human secreted protein sequence clone cn621_8.	994	94
889	Y41293	Homo sapiens	Human soluble protein ZTMPO-1.	4595	99
890	G03714	Homo sapiens	Human secreted protein, SEQ ID NO: 7795.	147	63
891	AF208856	Homo sapiens	BM-014	1012	99
892	U29195	Homo sapiens	neuronal pentraxin II	2002	98
893	X68149	Homo sapiens	Burkitt lymphoma receptor 1	1953	100
894	Y94914	Homo sapiens	Human secreted protein clone pw337_6 protein sequence SEQ ID NO:34.	537	100
895	W61630	Homo sapiens	Clone HNFGW06 of EGFR receptor family.	326	63
896	M24110	Homo sapiens	G0S19-2 peptide precursor	481	100
897	Z68747	Homo sapiens	imogen 38	2018	99
898	AF186112	Homo sapiens	neurokinin B-like protein ZNEUROK1	619	100
899	AF225420	Homo sapiens	AD025	734	100

SEQ	Accession	Species	Description	Smith-	T%
ID	No.	- Preside		Waterman	Identity
NO:				Score	
900	P60657	Homo sapiens	Sequence of human lipocortin.	1835	100
901	M27288	Homo sapiens	oncostatin M	1297	99
902	W85737	Homo sapiens	Polypeptide with transmembrane domain.	749	100
903	G01349	Homo sapiens	Human secreted protein, SEQ ID NO: 5430.	650	99
904	Y00261	Homo sapiens	Human secreted protein encoded by gene 4.	1133	99
905	AF039688	Homo sapiens	antigen NY-CO-3	771	99
906	AB007836	Homo sapiens	Hic-5	2544	100
907	AB017507	Homo sapiens	Apg12	224	100
908	AK000056	Homo sapiens	unnamed protein product	1537	98
909	Y86299	Homo sapiens	Human secreted protein HFOXB55, SEQ ID NO:214.	427	100
910	AF231023	Homo sapiens	protocadherin Flamingo 1	7393	99
911	Y14134	Homo sapiens	Vascular endothelial cell growth inhibitor beta protein sequence.	1319	100
912	Z90420	Homo sapiens	Human GDF-3 (hGDF-3) polypeptide encoding cDNA.	1950	100
913	Y19757	Homo sapiens	SEQ ID NO 475 from WO9922243.	1361	100
914	G03172	Homo sapiens	Human secreted protein, SEQ ID NO: 7253.	112	48
915	U14971	Homo sapiens	ribosomal protein S9	886	90
916	AF172854	Homo sapiens	cardiotrophin-like cytokine CLC	1204	99
917	AC005525	Homo sapiens	F22162_1	1963	100
918	AF166350	Homo sapiens	ST7 protein	4711	99
919	Y87285	Homo sapiens	Human signal peptide containing protein HSPP-62 SEQ ID NO:62.	430	100
920	Y36131	Homo sapiens	Human secreted protein #3.	465	88
921	AF193766	Homo sapiens	cytokine-like protein C17	724	100
922	Y95013	Homo sapiens	Human secreted protein vc48_1, SEQ ID NO:66.	357	100
923	X75208	Homo sapiens	protein tyrosine kinase-receptor	5256	100
924	Y96202	Homo sapiens	IkappaB kinase (IKK) binding protein, Y2H56.	813	98
925	AB039886	Homo sapiens	down-regulated in gastric cancer	785	78
926	G03368	Homo sapiens	Human secreted protein, SEQ ID NO: 7449.	55	50
927	Y48606	Homo sapiens	Human breast tumour-associated protein 67.	539	100
928	Y36151	Homo sapiens	Human secreted protein #23.	668	100
929	AF110399	Homo sapiens	elongation factor Ts	1666	100
930	AF210317	Homo sapiens	facilitative glucose transporter family member GLUT9	2763	99
931	Y73328	Homo sapiens	HTRM clone 082843 protein sequence.	931	100
932	G01959	Homo sapiens	Human secreted protein, SEQ ID NO: 6040.	274	100
933	U47924	Homo sapiens	B-cell receptor associated protein	1469	100
934	G03827	Homo sapiens	Human secreted protein, SEQ ID NO: 7908.	529	93
935	AB039371	Homo sapiens	mitochondrial ABC transporter 3	196	63
936	X56385	Canis	rab8	1064	100
937	B08906	familiaris Homo sapiens	Human secreted protein sequence encoded by	117	44
			gene 16 SEQ ID NO:63.		
938 939	M13692 Y53886	Homo sapiens Homo sapiens	A suppressor of cytokine signalling protein	1064 515	99 42
940	Y16630	Homo sapiens	designated HSCOP-6.  Human Putative Adrenomedullin Receptor (PAR).	1904	99
941	AC005102	Homo sapiens	small inducible cytokine subfamily A member 24	627	99
942	M12886	Homo sapiens	T-cell receptor beta chain	1289	81
943	AF226046	Homo sapiens	GK003	1049	98
944	Y36078	Homo sapiens	Extended human secreted protein sequence, SEQ ID NO. 463.	667	100
945	M22877	Homo sapiens	cytochrome c	565	100
946	W67869	Homo sapiens	Human secreted protein encoded by gene 63 clone HHGDB72.	551	93
947	W67859	Homo sapiens	Human secreted protein encoded by gene 53 clone HBMCL41.	283	100
948	W85726	Homo sapiens	Novel protein (Clone BG33_7).	789	100
949	AJ242015	Homo sapiens	eMDC II protein	4236	100
950	G04075	Homo sapiens	Human secreted protein, SEQ ID NO: 8156.	567	99

SEQ	Accession	Species	Description	Smith-	1%
ID	No.			Waterman	Identity
NO:				Score	
951	AF110645	Homo sapiens	candidate tumor suppressor p33 ING1 homolog	1314	100
952	Y36111	Homo sapiens	Extended human secreted protein sequence, SEQ	402	70
		1	ID NO. 496.		
953	AB012109	Homo sapiens	APC10	990	100
954	AF246221	Homo sapiens	transmembrane protein BRI	1405	100
955	AF054986	Homo sapiens	putative transmembrane GTPase	1883	100
956	W74726	Homo sapiens	Human secreted protein fg949_3.	1879	100
957	Y27096	Homo sapiens	Human viral receptor protein (ACVRP).	1581	100
958	AJ222967	Homo sapiens	cystinosin	1920	100
959	Y53052	Homo sapiens	Human secreted protein clone df202 3 protein	587	100
			sequence SEQ ID NO:110.		1
960	G02694	Homo sapiens	Human secreted protein, SEQ ID NO: 6775.	283	100
961	AF151855	Homo sapiens	CGI-97 protein	1214	96
962	U26592	Homo sapiens	diabetes mellitus type I autoantigen	250	65
963	AL050306	Homo sapiens	dJ475B7.2 (novel protein)	3796	100
964	AF078859	Homo sapiens	PTD004	2089	100
965	AB020315	Homo sapiens	homologue of mouse dkk-1 gene:Acc# AF030433	1466	100
966	X04571	Homo sapiens	precursor polypeptide (AA -22 to 1185)	6580	99
967	AF146019	Homo sapiens	hepatocellular carcinoma antigen gene 520	993	99
968	AF071002	Homo sapiens	minK-related peptide 1; MiRP1	632	100
969	AB021227	Homo sapiens	membrane-type-5 matrix metalloproteinase	3545	100
970	AF180920		cyclin L ania-6a	1579	100
970 971	AF180920 AF105365	Homo sapiens	K-Cl cotransporter KCC4	5621	99
		Homo sapiens	K-Ci cotransporter KCC4		
972	AF083248	Homo sapiens	ribosomal protein L26 homolog	739	100
973	AJ132429	Homo sapiens	hyperpolarization-activated cyclic nucleotide gated cation channel hHCN4	6295	100
974	W61619	Homo sapiens	Clone HTPEF86 of TM4SF superfamily.	454	100
975	AF155100	Homo sapiens	zinc finger protein NY-REN-21 antigen	2261	100
976	AF275948	Homo sapiens	ABCA1	11763	99
977	AB026891	Homo sapiens	cystine/glutamate transporter	2552	100
978	AF117657	Homo sapiens	thyroid hormone receptor-associated protein complex component TRAP80	3348	99
979	AF044201	Rattus .	neural membrane protein 35; NMP35	1570	92
		norvegicus			
980	AF119297	Homo sapiens	neuroendocrine-specific protein-like protein I	1170	99
981	AF155652	Homo sapiens	potassium channel modulatory factor	1983	99
982	W88499	Homo sapiens	Human stomach carcinoma clone HP10412- encoded protein.	1553	99
983	Z56281	Homo sapiens	interferon regulatory factor 3	2012	98
984	AB026125	Homo sapiens	ART-4	2160	100
985	Y14482	Homo sapiens	Fragment of human secreted protein encoded by gene 17.	172	70
986	AB023888	Homo sapiens	b-chemokine receptor CCR4	1895	100
987	W27291	Homo sapiens	Human H1075-1 secreted protein 5' end.	712	100
988	AF153450	Manduca	juvenile hormone esterase binding protein	226	32
000	002607	sexta	17	104	00
989	G03697	Homo sapiens	Human secreted protein, SEQ ID NO: 7778.	194	88
990	AF204159	Homo sapiens	potassium large conductance calcium-activated channel beta 3a subunit	1486	100
991	G02061	Homo sapiens	Human secreted protein, SEQ ID NO: 6142.	558	99
992	AL031266	Caenorhabditi s elegans	VM106R.1	327	40
993	Y66749	Homo sapiens	Membrane-bound protein PRO1124.	4730	99
994	G01246	Homo sapiens	Human secreted protein, SEQ ID NO: 5327.	141	77
995	AF133845	Homo sapiens	corin	5811	99
995	AF133843 AF117756	Homo sapiens	thyroid hormone receptor-associated protein	4999	100
00=	177.600.55	ļ.,	complex component TRAP150		100
997	W62066	Homo sapiens	Human stem cell antigen 2.	284	93
998	Y87173	Homo sapiens	Human secreted protein sequence SEQ ID NO:212.	725	100
999	Y13379	Homo sapiens	Amino acid sequence of protein PRO263.	1654	99
1000	Y95008	Homo sapiens	Human secreted protein vf3_1, SEQ ID NO:56.	676	47
		,	membrane associated protein SLP-2	1747	100

SEQ	Accession	Species	Description	Smith-	%
ID	No.			Waterman	Identity
NO:			, , , , , , , , , , , , , , , , , , ,	Score	
1002	G01234	Homo sapiens	Human secreted protein, SEQ ID NO: 5315.	398	96
1003	W73420	Homo sapiens	Human secreted protein encoded by Gene No. 24.	2150	100
1004	X12791	Homo sapiens	19kD SRP-protein (AA 1 - 144)	742	100
1005	M23323	Homo sapiens	membrane protein	642	100
1006	X63745	Homo sapiens	KDEL receptor	326	98
1007	Y35997	Homo sapiens	Extended human secreted protein sequence, SEQ ID NO. 382.	824	99
1008	AB032918	Hylobates moloch	dopamine receptor D4	92	35
1009	Y91680	Homo sapiens	Human secreted protein sequence encoded by gene 81 SEQ ID NO:353.	1372	99
1010	AL136125	Homo sapiens	dJ304B14.1 (novel protein)	825	98
1011	G03733	Homo sapiens	Human secreted protein, SEQ ID NO: 7814.	379	98
1012	Y17531	Homo sapiens	Human secreted protein clone BL205 14 protein.	818	97
1013	G00724	Homo sapiens	Human secreted protein, SEQ ID NO: 4805.	462	100
1014	AF288092	Naegleria gruberi	haem lyase	114	37
1015	AB045292	Homo sapiens	M83 protein	3867	99
1016	X15940	Homo sapiens	ribosomal protein L31 (AA 1-125)	644	100
1017	Y94873	Homo sapiens	Human protein clone HP02632.	1876	100
1018	AL024498	Homo sapiens	dJ417M14.1 (novel protein)	589	100
1019	X83425	Homo sapiens	Lutheran blood group glycoprotein	3054	99
1020	W03516	Homo sapiens	Prostaglandin DP receptor.	1864	100
1021	G03960	Homo sapiens	Human secreted protein, SEQ ID NO: 8041.	398	100
1022	Y91689	Homo sapiens	Human secreted protein sequence encoded by gene 93 SEQ ID NO:362.	768	100
1023	AE000660	Homo sapiens	hADV36S1	573	100
1024	AF132965	Homo sapiens	CGI-31 protein	1550	100
1025	W92380	Homo sapiens	Human TR-interacting protein S103a.	1466	97
1026	R66278	Homo sapiens	Therapeutic polypeptide from glioblastoma cell line.	830	100
1027	X65614	Homo sapiens	S100P calcium-binding protein	476	100
1028	Y41741	Homo sapiens	Human PRO704 protein sequence.	1323	100
1029	AJ001014	Homo sapiens	RAMP1	806	100
1030	W63682	Homo sapiens	Human secreted protein 2.	1354	99
1031	AK023007	Homo sapiens	unnamed protein product	766	100
1032	W97900	Homo sapiens	Human SR-BI class B scavenger.	2672	99
1033	Y82453	Homo sapiens	Human TGC-440 secretory protein SEQ ID	639	99
			NO:1.		
1034	Y73473	Homo sapiens	Human secreted protein clone yd178_1 protein sequence SEQ ID NO:168.	752	93
1035	Y86468	Homo sapiens	Human gene 48-encoded protein fragment, SEQ ID NO:383.	96	90
1036	U09813	Homo sapiens	mitochondrial ATP synthase subunit 9 precursor	698	100
1037	AJ242832	Homo sapiens	calpain	3699	99
1038	X66403	Homo sapiens	acetylcholine receptor epsilon subunit CHRNE	2574	100
1039	AJ242730	Homo sapiens	polyhomeotic 2	1310	100
1040	AF169968	Mus musculus	DNA binding protein DESRT	1453	80
1041	X52563	Bos taurus	permability increasing protein	383	29
1042	G00368	Homo sapiens	Human secreted protein, SEQ ID NO: 4449.	75	50
1043	G02532	Homo sapiens	Human secreted protein, SEQ ID NO: 6613.	60	53
1044	M94582	Homo sapiens	interleukin 8 receptor B	1850	100
1045	AL080239	Homo sapiens	bG256O22.1 (similar to IGFALS (insulin-like growth factor binding protein, acid labile subunit))	1704	50
1046	AF125101	Homo sapiens	HSPC040 protein	580	100
1047	W74809	Homo sapiens	Human secreted protein encoded by gene 81 clone HMWDN32.	176	100
1048	AL022238	Homo sapiens	dJ1042K10.4 (novel protein)	2201	100
1049	W88667	Homo sapiens	Secreted protein encoded by gene 134 clone HAIBP89.	1559	99
			14 14 14 14 14 14 14 14 14 14 14 14 14 1		1

1052 Y21 1053 AL1 1054 Y76 1055 AJ2 1056 Y27 1057 D14 1058 AF1 1059 AL6 1060 AF2 1061 Y27 1062 Z11 1063 AF1 1064 AF1 1065 Y41 1066 AJ2 1067 Y36 1069 Y94 1070 W66 1071 X03 1072 AL6 1073 X82 1074 G03 1075 Y36 1076 G03 1077 L25 1078 Y91 1079 G01 1079 G01 1080 AB6 1081 AB6 1082 L13 1083 W73 1084 G03 1085 G04 1086 AF0 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W83	721851 AL163815 776200 AJ276567 727620 014530 AF132000 AL031778	<del></del>		Score	Identity
1053 AL1 1054 Y76 1055 AJ2 1056 Y27 1057 D14 1058 AF1 1059 AL6 1060 AF2 1061 Y27 1062 Z11 1063 AF1 1064 AF1 1065 Y41 1066 AJ2 1067 Y36 1069 Y94 1070 W66 1071 X03 1072 AL6 1071 X03 1072 AL6 1073 X82 1076 G03 1077 L25 1078 Y91 1079 G01 1080 AB6 1081 AB6 1082 L13 1083 W7:	AL163815 776200 AJ276567 727620 D14530 AF132000	Homo sapie	Fragment of human secreted protein encoded by gene 81.	1318	98
1054 Y76 1055 AJ2 1056 Y27 1057 D14 1058 AF1 1059 ALC 1060 AF2 1061 Y27 1062 Z11 1063 AF1 1064 AF1 1065 Y41 1066 AJ2 1067 Y36 1068 Y94 1069 Y94 1070 W66 1071 X03 1072 ALC 1073 X82 1074 G03 1075 Y36 1076 G03 1077 L25 1078 Y91 1079 G01 1080 ABC 1081 ABC 1082 L13 1083 W73 1084 G03 1085 G04 1086 AFC 1087 G00 1088 G04 1088 G04 1088 G04 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W83	776200 AJ276567 727620 D14530 AF132000		Human signal peptide-contianing protein (SIGP) (clone ID 2328134).	1643	95
1055         AJ2           1056         Y27           1057         D14           1058         AF1           1059         ALG           1060         AF2           1061         Y27           1062         Z11           1063         AF1           1064         AF1           1065         Y41           1066         AJ2           1067         Y36           1068         Y94           1070         W6           1071         X03           1072         ALG           1073         X82           1074         G03           1075         Y36           1076         G03           1077         L25           1078         Y91           1079         G01           1080         AB6           1081         AB6           1082         L13           1083         W7           1084         G03           1085         G04           1086         AF0           1087         G00           1088         G04	AJ276567 (27620 014530 AF132000	63815 Arabidopsis thaliana	putative protein	661	62
1056         Y27           1057         D14           1058         AF1           1059         ALG           1060         AF2           1061         Y27           1062         Z11           1063         AF1           1064         AF1           1065         Y41           1066         AJ2           1067         Y36           1068         Y94           1070         W6           1071         X03           1072         ALG           1073         X82           1074         G03           1075         Y36           1076         G03           1077         L25           1078         Y91           1079         G01           1080         AB6           1081         AB6           1082         L13           1083         W7           1084         G03           1085         G04           1087         G00           1088         G04           1091         S72           1092         W8	(27620 014530 AF132000	200 Homo sapie	Human secreted protein encoded by gene 77.	262	100
1057         D14           1058         AFI           1059         ALG           1059         ALG           1059         ALG           1060         AFI           1061         Y27           1062         Z11           1063         AFI           1064         AFI           1065         Y41           1066         AJ2           1067         Y36           1070         W6           1071         X03           1072         ALG           1073         X82           1074         G03           1075         Y36           G03         1077         L25           1078         Y91           1079         G01           1080         ABG           1081         ABG           1082         L13           1083         W7:           1084         G03           1085         G04           1087         G00           1088         G04           1090         G04           1091         S72           1092         <	014530 AF132000			1160	100
1058         AF1           1059         AL6           1059         AL6           1059         AL6           1061         Y27           1062         Z11           1063         AF1           1065         Y41           1065         Y41           1066         AJ2           1067         Y36           1068         Y94           1070         W6           1071         X03           1072         AL6           1073         X82           1074         G03           1075         Y36           1076         G03           1077         L25           1078         Y91           1079         G01           1080         AB6           1081         AB6           1082         L13           1083         W7:           1084         G03           1085         G04           1087         G00           1088         G04           1090         G04           1091         S72           1092         W8:	AF132000			154	96
1059 ALG 1060 AF2 1061 Y27 1062 Z11 1063 AF1 1064 AF1 1065 Y41 1066 AJ2 1067 Y36 1068 Y94 1069 Y94 1070 W6 1071 X03 1072 ALG 1073 X82 1074 G03 1075 Y36 1076 G03 1077 L25 1078 Y91 1079 G01 1080 ABG 1081 ABG 1081 ABG 1082 L13 1083 W7: 1084 G03 1085 G04 1086 AFG 1087 G00 1088 G04 1088 G04 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88				745	100
1060 AF2 1061 Y27 1062 Z11 1063 AF1 1064 AF1 1065 Y41 1066 AJ2 1067 Y36 1068 Y94 1069 Y94 1070 W6 1071 X03 1072 ALC 1073 X82 1074 G03 1075 Y36 1076 G03 1077 L25 1078 Y91 1079 G01 1080 ABC 1081 ABC 1081 ABC 1082 L13 1083 W7: 1084 G03 1085 G04 1086 AFC 1087 G00 1088 G04 1088 G04 1088 G04 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	AL031778		TADA1 protein	1132	100
1061         Y27           1062         Z11           1063         AF1           1064         AF1           1065         Y41           1066         AJ2           1067         Y36           1068         Y94           1069         Y94           1070         W6-           1071         X03           1072         ALC           1073         X82           1074         G03           1075         Y36           1076         G03           1077         L25           1078         Y91           1080         ABC           1081         ABC           1082         L13           1083         W7:           1084         G03           1085         G04           1086         AFC           1087         G00           1088         G04           1090         G04           1091         S72           1092         W88			receptor (peripheral) (MBR, PBR, PBKS, IBP, Isoquinoline-binding protein)) LIKE protein)	920	100
1062         Z11           1063         AF1           1064         AF1           1065         Y41           1066         AJ2           1067         Y36           1068         Y94           1069         Y94           1070         W6-           1071         X03           1072         AL0           1073         X82           1074         G03           1075         Y36           1076         G03           1077         L25           1078         Y91           1080         AB0           1081         AB0           1082         L13           1083         W7:           1084         G03           1085         G04           1086         AF0           1087         G00           1088         G04           1090         G04           1091         S72           1092         W88           1093         W8:	AF227135			134	33
1063 AF1 1064 AF1 1065 Y41 1066 AJ2 1067 Y36 1068 Y94 1069 Y94 1070 W66 1071 X03 1072 ALC 1073 X82 1074 G03 1077 L25 1076 G03 1077 L25 1078 Y91 1080 ABC 1081 ABC 1082 L13 1083 W7: 1084 G03 1085 G04 1086 AFC 1087 G00 1088 G04 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	(27575	575 Homo sapie		1392	100
1064 AF1 1065 Y41 1066 AJ2 1067 Y36 1068 Y94 1069 Y94 1070 W66 1071 X03 1072 ALC 1073 X82 1074 G03 1077 L25 1078 Y91 1079 G01 1080 ABC 1081 ABC 1082 L13 1083 W7: 1084 G03 1085 G04 1086 AFC 1087 G00 1088 G04 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	Z11697			1088	100
1065 Y41 1066 AJ2 1067 Y36 1068 Y94 1069 Y94 1070 W66 1071 X03 1072 ALC 1073 X82 1074 G03 1075 Y36 1076 G03 1077 L25 1078 Y91 1080 ABC 1081 ABC 1081 ABC 1082 L13 1083 W7: 1084 G03 1085 G04 1087 G00 1088 G04 1088 AFC 1089 AFI 1090 G04 1091 S72 1092 W88	AF123757			819	100
1066 AJ2 1067 Y36 1068 Y94 1069 Y94 1070 W66 1071 X03 1072 ALC 1073 X82 1074 G03 1075 Y36 1076 G03 1077 L25 1078 Y91 1080 ABC 1081 ABC 1081 ABC 1082 L13 1083 W73 1084 G03 1085 G04 1086 AFC 1087 G00 1088 G04 1088 G04 1089 AFI 1090 G04 1091 S72 1092 W88	AF155135			2932	99
1067 Y36 1068 Y94 1069 Y94 1070 W66 1071 X03 1072 ALC 1073 X82 1074 G03 1077 L25 1078 Y91 1079 G01 1080 ABC 1081 ABC 1081 ABC 1082 L13 1083 W73 1084 G03 1085 G04 1086 AFC 1087 G00 1088 G04 1088 G04 1089 AFI 1090 G04 1091 S72 1092 W88	741674			936	99
1068 Y94 1069 Y94 1070 W66 1071 X03 1072 ALC 1073 X82 1074 G03 1075 Y36 1076 G03 1077 L25 1078 Y91 1080 ABC 1081 ABC 1082 L13 1083 W73 1084 G03 1085 G04 1086 AFC 1087 G00 1088 G04 1089 AFI 1090 G04 1091 S72 1092 W88	AJ250042			2575	100
1069 Y94 1070 W6 1071 X03 1072 ALC 1073 X82 1074 G03 1075 Y36 1076 G03 1077 L25 1078 Y91 1079 G01 1080 ABC 1081 ABC 1082 L13 1083 W7: 1084 G03 1085 G04 1086 AFC 1087 G00 1088 G04 1088 G04 1088 G04 1089 AFI 1090 G04 1091 S72 1092 W88	736087		ID NO. 472.	770	85
1070 W64 1071 X03 1072 ALC 1073 X82 1074 G03 1075 Y36 1076 G03 1077 L25 1078 Y91 1079 G01 1080 ABG 1081 ABG 1082 L13 1083 W73 1084 G03 1085 G04 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	794959	•	sequence SEQ ID NO:124.	301	100
1071 X03 1072 ALC 1073 X82 1074 G03 1075 Y36 1076 G03 1077 L25 1078 Y91 1079 G01 1080 ABG 1081 ABG 1082 L13 1083 W73 1084 G03 1085 G04 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W83	794959	•	sequence SEQ ID NO:124.	301	100
1072 ALC 1073 X82 1074 G03 1075 Y36 1076 G03 1077 L25 1078 Y91 1079 G01 1080 ABC 1081 ABC 1082 L13 1083 W73 1084 G03 1085 G04 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	V64535			2014	99
1073   X82   1074   G03   1075   Y36   1076   G03   1077   L25   1078   Y91   1079   G01   1080   ABG   1081   ABG   1082   L13   1084   G03   1085   G04   1086   AFG   1086   AFG   1087   G00   G04   1090   G04   1090   G04   1091   S72   1092   W88   1093   W85   1094   1095   W85   1097   W85   W	(03145			148	50
1074 G03 1075 Y36 1076 G03 1077 L25 1078 Y91 1079 G01 1080 AB0 1081 AB0 1082 L13 1083 W7: 1084 G03 1085 G04 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	L031177			821	91
1075 Y36 1076 G03 1077 L25 1078 Y91 1079 G01 1080 AB0 1081 AB0 1082 L13 1083 W7: 1084 G03 1085 G04 1086 AF0 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88				249	62
1076 G03 1077 L25 1078 Y91 1079 G01 1080 AB0 1081 AB0 1082 L13 1083 W7: 1084 G03 1085 G04 1086 AF0 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	303213			99	47
1077 L25 1078 Y91 1079 G01 1080 AB0 1081 AB0 1082 L13 1083 W7: 1084 G03 1085 G04 1086 AF0 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	736233			506 424	55
1078 Y91 1079 G01 1080 AB0 1081 AB0 1082 L13 1083 W7: 1084 G03 1085 G04 1086 AF0 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88			Human secreted protein, SEQ ID NO: 7268. ribosomal protein L10	332	98
1079 G01 1080 AB0 1081 AB0 1082 L13 1083 W73 1084 G03 1085 G04 1086 AF0 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	123899 (91447			898	76 97
1080 AB0 1081 AB0 1082 L13 1083 W7: 1084 G03 1085 G04 1086 AF0 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88			gene 48 SEQ ID NO:168.		
1081 ABG 1082 L13 1083 W7: 1084 G03 1085 G04 1086 AFG 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88 1093 W8:	AB039723		Human secreted protein, SEQ ID NO: 5943. WNT receptor frizzled-3	290 1376	89 92
1082 L13 1083 W7: 1084 G03 1085 G04 1086 AFC 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	AB039723 AB020527			269	100
1083 W7: 1084 G03 1085 G04 1086 AFC 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88				499	80
1084 G03 1085 G04 1086 AFG 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	W75098	l *		143	81
1085 G04 1086 AFG 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88		•	clone HSXBI25.		
1086 AFC 1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	303564			83	51
1087 G00 1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	304063			88	43
1088 G04 1089 AF1 1090 G04 1091 S72 1092 W88	F090942			124	64
1089 AF1 1090 G04 1091 S72 1092 W88				129 126	41
1090 G04 1091 S72 1092 W88 1093 W88	AF140631			364	36 82
1091 S72 1092 W88 1093 W83	304063			114	32
1092 W88 1093 W83	72304		LMW G-protein	146 .	83
	V88708			405	100
	V85612	612 Homo sapie		4358	97
	753012			1013	99
1095 Y92	792345	345 Homo sapie		409	100
1096 AF0		90942 Homo sapie		147	60
	F090942			166	58
1098 X56	F090942 24521			490	70
			Human secreted protein, SEQ ID NO: 8144.	83	35
1100 Y02	24521			149	59

ID	Accession	Species	Description	Smith-	%
TATO.	No.			Waterman	Identity
NO: 1101	AF119851	Homo sapiens	PD 01500	Score 183	72
1101	G04086		PRO1722	207	
1102	G04086	Homo sapiens	Human secreted protein, SEQ ID NO: 8167.	91	62 52
	X74856	Homo sapiens	Human secreted protein, SEQ ID NO: 8144.		
1104		Mus musculus	ribosomal protein L28	128	69
1105	G03789	Homo sapiens	Human secreted protein, SEQ ID NO: 7870.	130	62
106	G03133	Homo sapiens	Human secreted protein, SEQ ID NO: 7214.	122	48
107	G03040	Homo sapiens	Human secreted protein, SEQ ID NO: 7121.	69	43
108	AF039942	Homo sapiens	HCF-binding transcription factor Zhangfei	744	99
1109	AF201951	Homo sapiens	high affinity immunoglobulin epsilon receptor beta subunit	738	94
1110	AF111108	Mus musculus	transient receptor potential 2	223	79
1111	AF119900	Homo sapiens	PRO2822	144	59
1112	Y16589	Homo sapiens	A protein that interacts with presentlins.	265	39
113	G02872	Homo sapiens	Human secreted protein, SEQ ID NO: 6953.	178	67
1114	Y02999	Homo sapiens	Fragment of human secreted protein encoded by gene 121.	164	63
115	Y30811	Homo sapiens	Human secreted protein encoded from gene 1.	1217	99
116	X51394	Xenopus laevis	APEG precursor protein	130	40
117	M27826	Homo sapiens	neutral protease large subunit	442	65
118	G03371	Homo sapiens	Human secreted protein, SEQ ID NO: 7452.	72	60
119	G03602	Homo sapiens	Human secreted protein, SEQ ID NO: 7683.	491	97
120	Y35906	Homo sapiens	Extended human secreted protein sequence, SEQ ID NO. 155.	244	97
121	G03714	Homo sapiens	Human secreted protein, SEQ ID NO: 7795.	122	65
122	Y00337	Homo sapiens	Human secreted protein encoded by gene 81.	110	90
123	AF084830	Homo sapiens	two pore domain K+ channel; TASK-2	703	94
124	AF212862	Homo sapiens	membrane interacting protein of RGS16	442	88
1125	W64469	Homo sapiens	Human secreted protein from clone CW795 2.	191	53
1126	G01361	Homo sapiens	Human secreted protein, SEQ ID NO: 5442.	154	100
1127	G01361	Homo sapiens	Human secreted protein, SEQ ID NO: 5442.	165	100
1128	Y84320	Homo sapiens	Human cardiovascular system associated protein kinase-1.	815	99
1129	G02105	Homo sapiens	Human secreted protein, SEQ ID NO: 6186.	88	73
1130	Y32923	Homo sapiens	Transmembrane domain containing protein clone HP01512.	700	100
1131	Y29817	Homo sapiens	Human synapse related glycoprotein 2.	260	91
1132	Y91644	Homo sapiens	Human secreted protein sequence encoded by gene 43 SEQ ID NO:317.	525	96
1133	Y91449	Homo sapiens	Human secreted protein sequence encoded by gene 49 SEQ ID NO:170.	542	100
1134	AB017908	Homo sapiens	4F2 light chain	2399	93
1135	X51760	Homo sapiens	zinc finger protein (583 AA)	312	55
1136	Y99426	Homo sapiens	Human PRO1604 (UNQ785) amino acid sequence SEQ ID NO:308.	917	72
1137	G03790	Homo sapiens	Human secreted protein, SEQ ID NO: 7871.	102	50
	AF155106	Homo sapiens	NY-REN-36 antigen	768	91
	1 100		dJ28H20.1 (novel protein similar to membrane	117	50
1138	AL031055	Homo sapiens		117	1
1138	Ì		transport proteins)	_	
1138 1139 1140	AF011359	Bos taurus	transport proteins) regulator of G-protein signaling 7	138	96
138 139 140 141	AF011359 Y70018	Bos taurus Homo sapiens	transport proteins) regulator of G-protein signaling 7 Human Protease and associated protein-12 (PPRG-12).	138 623	96 100
138 139 140 141	AF011359 Y70018 G04091	Bos taurus Homo sapiens Homo sapiens	transport proteins) regulator of G-protein signaling 7 Human Protease and associated protein-12 (PPRG-12). Human secreted protein, SEQ ID NO: 8172.	138 623	96 100 38
1138 1139 1140 1141 1142 1143	AF011359 Y70018 G04091 AB030235	Bos taurus Homo sapiens Homo sapiens Canis familiaris	transport proteins) regulator of G-protein signaling 7 Human Protease and associated protein-12 (PPRG-12). Human secreted protein, SEQ ID NO: 8172. D4 dopamine receptor	138 623 113 89	96 100 38 48
1138 1139 1140 1141 1142 1143	AF011359 Y70018 G04091 AB030235 Y94922	Bos taurus Homo sapiens Homo sapiens Canis familiaris Homo sapiens	transport proteins) regulator of G-protein signaling 7 Human Protease and associated protein-12 (PPRG-12). Human secreted protein, SEQ ID NO: 8172. D4 dopamine receptor  Human secreted protein clone pv6_1 protein sequence SEQ ID NO:50.	138 623 113 89 539	96 100 38 48
1138 1139 1140 1141 1142 1143 1144	AF011359 Y70018 G04091 AB030235 Y94922 X99962	Bos taurus Homo sapiens Homo sapiens Canis familiaris Homo sapiens Homo sapiens	transport proteins) regulator of G-protein signaling 7 Human Protease and associated protein-12 (PPRG-12). Human secreted protein, SEQ ID NO: 8172. D4 dopamine receptor  Human secreted protein clone pv6_1 protein sequence SEQ ID NO:50. rab-related GTP-binding protein	138 623 113 89 539	96 100 38 48 88
1138 1139 1140 1141 1142 1143 1144 1145 1146	AF011359 Y70018 G04091 AB030235 Y94922 X99962 G03807	Bos taurus Homo sapiens Canis familiaris Homo sapiens Homo sapiens Homo sapiens	transport proteins) regulator of G-protein signaling 7 Human Protease and associated protein-12 (PPRG-12). Human secreted protein, SEQ ID NO: 8172. D4 dopamine receptor  Human secreted protein clone pv6_1 protein sequence SEQ ID NO:50. rab-related GTP-binding protein Human secreted protein, SEQ ID NO: 7888.	138 623 113 89 539 398 168	96 100 38 48 88 96 79
1138 1139 1140 1141 1142 1143 1144	AF011359 Y70018 G04091 AB030235 Y94922 X99962	Bos taurus Homo sapiens Homo sapiens Canis familiaris Homo sapiens Homo sapiens	transport proteins) regulator of G-protein signaling 7 Human Protease and associated protein-12 (PPRG-12). Human secreted protein, SEQ ID NO: 8172. D4 dopamine receptor  Human secreted protein clone pv6_1 protein sequence SEQ ID NO:50. rab-related GTP-binding protein	138 623 113 89 539	96 100 38 48 88

SEQ ID NO:	Accession No.	Species	Description	Smith- Waterman Score	% Identity
		s elegans	cerevisiae zinc resistance protein		
1150	G03438	Homo sapiens	Human secreted protein, SEQ ID NO: 7519.	117	62
1151	G01003	Homo sapiens	Human secreted protein, SEQ ID NO: 5084.	181,	80
1152	G03798	Homo sapiens	Human secreted protein, SEQ ID NO: 7879.	198	63
1153	X88799	Oryza sativa	DNA binding protein	95	41
1154	D85245	Homo sapiens	TR3beta	155	96
1155	R74272	Homo sapiens	Tumour suppressor protein, p53.	341	87
1156	Y86265	Homo sapiens	Human secreted protein HUSXE77, SEQ ID NO:180.	99	41
1157·	G02577	Homo sapiens	Human secreted protein, SEQ ID NO: 6658.	263	98
1158	AF104334	Homo sapiens	putative organic anion transporter	185	42
1159	G01393	Homo sapiens	Human secreted protein, SEQ ID NO: 5474.	173	57
1160	W75771	Homo sapiens	Human GTP binding protein APD08.	224	81
1161	AF216833	Homo sapiens	M-ABC2 protein	410	83
1162	W67816	Homo sapiens	Human secreted protein encoded by gene 10 clone HCEMU42.	1156	100
1163	AF119851	Homo sapiens	PRO1722	230	70
1164	Y87252	Homo sapiens	Human signal peptide containing protein HSPP- 29 SEQ ID NO:29.	113	31
1165	W64537	Homo sapiens	Human liver cell clone HP01148 protein.	338	82
1166	AF269286	Homo sapiens	HC6	134	64
1167	Y14482	Homo sapiens	Fragment of human secreted protein encoded by gene 17.	149	51
1168	D90789	Escherichia coli	Dipeptide transport system permease protein DppC.	411	90
1169	R63783	Homo sapiens	TG0847 protein.	344	90
1170	Y45274	Homo sapiens	Human secreted protein encoded from gene 18.	478	98
1171	D64154	Homo sapiens	Mr 110,000 antigen	347	96
1172	AB026256	Homo sapiens	organic anion transporter OATP-B	311	67
1173	G00357	Homo sapiens	Human secreted protein, SEQ ID NO: 4438.	60	52
1174	D87717	Homo sapiens	similar to human GTPase-activating protein(A49869)	178	59
1175	M64716	Homo sapiens	ribosomal protein	391	78
1176	R08330	Homo sapiens	Human IL-7 receptor clone H6.	285	67
1177	L06505	Homo sapiens	ribosomal protein L12	242	72
1178	AJ251885	Homo sapiens	organic cation transporter (OCT2)	276	88
1179	G03258	Homo sapiens	Human secreted protein, SEQ ID NO: 7339.	155	71
1180	G01207	Homo sapiens	Human secreted protein, SEQ ID NO: 5288.	282	90
1181	AF181856	Rattus	tRNA selenocysteine associated protein	249	62
	12 101000	norvegicus	are a resistant appointed proton	1	02
1182	AF161524	Homo sapiens	HSPC176	138	90
1183	G03789	Homo sapiens	Human secreted protein, SEQ ID NO: 7870.	282	66
1184	Y02671	Homo sapiens	Human secreted protein encoded by gene 22 clone HMSJW18.	107	71
1185	G03797	Homo sapiens	Human secreted protein, SEQ ID NO: 7878.	88	69
1186	G03564	Homo sapiens	Human secreted protein, SEQ ID NO: 7645.	118	46
1187	AB032905	Hylobates concolor	dopamine receptor D4	96	37
1188	G00956	Homo sapiens	Human secreted protein, SEQ ID NO: 5037.	292	78
1189	G03258	Homo sapiens	Human secreted protein, SEQ ID NO: 7339.	178	79
1190	G03361	Homo sapiens	Human secreted protein, SEQ ID NO: 7442.	324	76
1191	AF117755	Homo sapiens	thyroid hormone receptor-associated protein complex component TRAP230	187	70
1192	Y70455	Homo sapiens	Human membrane channel protein-5 (MECHP-5).	202	67
1193	G03052	Homo sapiens	Human secreted protein, SEQ ID NO: 7133.	99	42
1194	G02607	Homo sapiens	Human secreted protein, SEQ ID NO: 6688.	192	76
1195	W29661	Homo sapiens	Homo sapiens CI542 2 clone secreted protein.	2001	98
1196	Y14104	Homo sapiens	Human GABAB receptor 1d protein sequence.	239	69
1197	X61972	Homo sapiens	macropain subunit iota	149	90
1198	G00534	Homo sapiens	Human secreted protein, SEQ ID NO: 4615.	145	51
1199	Y86260	Homo sapiens	Human secreted protein HELHN47, SEQ ID NO:175.	1089	89
	G02607	Homo sapiens	Human secreted protein, SEQ ID NO: 6688.	154	57

SEQ ID	Accession No.	Species	Description	Smith- Waterman	% Identity
NO:	1			Score	
1201	G00838	Homo sapiens	Human secreted protein, SEQ ID NO: 4919.	404	50
1202	M27826	Homo sapiens	neutral protease large subunit	202	49
1203	Y73424	Homo sapiens	Human secreted protein clone yi4_1 protein sequence SEQ ID NO:70.	265	61
1204	AF264014	Homo sapiens	scavenger receptor cysteine-rich type 1 protein M160 precursor	625	98
1205	Y36203	Homo sapiens	Human secreted protein #75.	219	59
1206	U78111	Gallus gallus	AQ	205	57
1207	AF095448	Homo sapiens	putative G protein-coupled receptor	416	76
1208	AF116715	Homo sapiens	PRO2829	127	75
1209	AF099137	Homo sapiens	MaxiK channel beta 2 subunit	475	95
1210	AF205718	Homo sapiens	hepatocellular carcinoma-related putative tumor suppressor	423	79
1211	Y27868	Homo sapiens	Human secreted protein encoded by gene No. 107.	224	70
1212	G00719	Homo sapiens	Human secreted protein, SEQ ID NO: 4800.	117	44
1213	G01009	Homo sapiens	Human secreted protein, SEQ ID NO: 5090.	351	73
1214	AF090942	Homo sapiens	PRO0657	124	70
1215	Y14427	Homo sapiens	Human secreted protein encoded by gene 17 clone HSIEA14.	99	77
1216	G03905	Homo sapiens	Human secreted protein, SEQ ID NO: 7986.	173	57
1217	Y57897	Homo sapiens	Human transmembrane protein HTMPN-21.	1173	100
1218	J00194	Homo sapiens	hla-dr antigen alpha chain	454	78
1219	Y59709	Homo sapiens	Secreted protein 76-28-3-A12-FL1.	470	92
1220	W81576	Homo sapiens	EBV-induced G-protein coupled receptor (EBI-	725	100
1221	W96745	Homo sapiens	polypeptide.     High affinity immunoglobulin E receptor-like	650	98
1222	Y35911	Homo sapiens	protein (IGERB).  Extended human secreted protein sequence, SEQ	135	31
1223	Y00278	Homo sapiens	ID NO. 160.  Human secreted protein encoded by gene 21.	260	95
1224	AF161422	Homo sapiens	HSPC304	568	90
1225	U14970	Homo sapiens	ribosomal protein S5	202	95
1226	G01733	Homo sapiens	Human secreted protein, SEQ ID NO: 5814.	610	100
1227	AF099973	Mus musculus	schlafen2	333	56
1228	G01218	Homo sapiens	Human secreted protein, SEQ ID NO: 5299.	155	81
1229	AF217188	Mus musculus	YIPIB	801	63
1230	AF176813	Homo sapiens	soluble adenylyl cyclase	275	100
1231	X98333	Homo sapiens	organic cation transporter	1704	100
1232	W74955	Homo sapiens	Human secreted protein encoded by gene 77 clone HOEAS24.	212	53
1233	Y94940	Homo sapiens	Human secreted protein clone yi62_1 protein sequence SEQ ID NO:86.	526	100
1234	U76618	Mus musculus	N-RAP	482	82
1235	AF044924	Homo sapiens	hook2 protein	380	97
1236	G01459	Homo sapiens	Human secreted protein, SEQ ID NO: 5540.	417	100
1237	AF000018	Homo sapiens	adapter protein	164	84
1238	W88633	Homo sapiens	Secreted protein encoded by gene 100 clone HE8EU04.	250	90
1239	W29660	Homo sapiens	Homo sapiens CH27_1 clone secreted protein.	697	98
1240	AF004161	Oryctolagus cuniculus	peroxisomal Ca-dependent solute carrier	154	52
1241	Y92710	Homo sapiens	Human membrane-associated protein Zsig24.	709	97
1242	Y95002	Homo sapiens	Human secreted protein vc34_1, SEQ ID NO:44.	908	88
1243	Y44905	Homo sapiens	Human potassium channel molecule ERG-LP2 partial protein.	325	100
1244	AF284422	Homo sapiens	cation-chloride cotransporter-interacting protein	511	97
1245	Y53629	Homo sapiens	A bone marrow secreted protein designated BMS115.	1888	93
1246	AB039371	Homo sapiens	mitochondrial ABC transporter 3	389	97
	Y35911	Homo sapiens	Extended human secreted protein sequence, SEQ	168	39

SEQ	Accession	Species	Description	Smith-	%
ID	No.			Waterman	Identity
NO:	ļ	<del></del>		Score	J
10.40	A E 0 5 0 0	1 B	ID NO. 160.	1	
1248	AF072509	Rattus norvegicus	glutamate receptor interacting protein 2	559	90
1249	AF247042	Homo sapiens	tandem pore domain potassium channel TRAAK	661	98
1250	B08974	Homo sapiens	Human secreted protein sequence encoded by	1087	97
		1	gene 27 SEQ ID NO:131.	1	1
1251	L15313	Caenorhabditi s elegans	putative	858	59
1252	Y29338	Homo sapiens	Human secreted protein clone it217_2 alternate reading frame protein.	278	75
1253	W01730	Homo sapiens	Human G-protein receptor HPRAJ70.	211	92
1254	G03074	Homo sapiens	Human secreted protein, SEQ ID NO: 7155.	294	83
1255	G01818	Homo sapiens	Human secreted protein, SEQ ID NO: 5899.	253	91
1256	AF286368	Homo sapiens	eppin-1	222	54
1257	AF220264	Homo sapiens	MOST-1	87	93
1258					
	G02227	Homo sapiens	Human secreted protein, SEQ ID NO: 6308.	281	78
1259	Y07970	Homo sapiens	Human secreted protein fragment #2 encoded from gene 26.	81	94
1260	R95332	Homo sapiens	Tumor necrosis factor receptor 1 death domain ligand (clone 3TW).	986	100
1261	AF140674	Homo sapiens	zinc metalloprotease ADAMTS6	172	36
1262	U28369	Homo sapiens	semaphorin V	237	67
1263	Y07049	Homo sapiens	Renal cancer associated antigen precursor sequence.	288	71
1264	Y36153	Homo sapiens	Human secreted protein #25.	187	80
1265	Y78114	Homo sapiens	Human cytokine signal regulator CKSR-2 SEQ ID NO:2.	723	93
1266	Y13397	TT		101	100
	AF030558	Homo sapiens	Amino acid sequence of protein PRO334.	191	100
1267	1	Rattus norvegicus	phosphatidylinositol 5-phosphate 4-kinase gamma	859	95
1268	U73167	Homo sapiens	candidate tumor suppressor gene LUCA-1	159	96
1269	AF190664	Mus musculus	LMBR2	552	76
1270	AL050332	Homo sapiens	dJ570F3.1 (homolog of the rat synaptic ras GTPase-activating protein p135 SynGAP)	820	98
1271	G02126	Homo sapiens	Human secreted protein, SEQ ID NO: 6207.	131	95
1272	AF125533	Homo sapiens	NADH-cytochrome b5 reductase isoform	253	92
1273	AL035661	Homo sapiens	dJ568C11.3 (novel AMP-binding enzyme similar to acetyl-coenzyme A synthethase (acetate-coA ligase))	1280	100
1274	AF064748	Mus musculus	S3-12	3523	61
1275	D17554	Homo sapiens	TAXREB107	377	78
1276	Y30715	Homo sapiens	Amino acid sequence of a human secreted	(10	1-22
			protein.	643	90
1277	AF146760	Homo sapiens	septin 2-like cell division control protein	707	100
1278	Y05069	Homo sapiens	Human PIGR-2 protein sequence.	281	46
1279	X59668	Oryctolagus cuniculus	aorta CNG channel (rACNG)	267	85
1280	G01051	Homo sapiens	Human secreted protein, SEQ ID NO: 5132.	489	98
1281	G03411	Homo sapiens	Human secreted protein, SEQ ID NO: 7492.	120	43
1282	AF055084	Homo sapiens	very large G-protein coupled receptor-1	1635	100
1283	AF117814	Mus musculus	odd-skipped related 1 protein	357	98
1284	U87318	Xenopus	NaDC-2	535	60
1285	AF061346	laevis Mus	Edp1 protein	452	68
1286	AB030182	musculus Mus	contains transmembrane (TM) region	582	68
1287	A13595	musculus synthetic	immunosuppresive protein PP15	185	97
		construct			1
1288	AF254411	Homo sapiens	ser/arg-rich pre-mRNA splicing factor SR-A1	837	100
1289	AF084205	Rattus	serine/threonine protein kinase TAO1	319	98
	1	norvegicus		1	1

SEQ	Accession	Species	Description	Smith-	%
ID NO:	No.	}		Waterman Score	Identity
1290	AF038563	Homo sapiens	mombrons are deted assessible lines 2	523	100
1290	AF034837	Homo sapiens	membrane associated guanylate kinase 2 double-stranded RNA specific adenosine	468	100
1291	AF034837	Homo sapiens	deaminase	468	100
1292	M15888	Bos taurus	endozepine-related protein precursor	937	87
1293	AB010692	Arabidopsis	ATP-dependent RNA helicase-like protein	636	45
1204	A T200002	thaliana		1500	100
1294 1295	AF209923 W67828	Homo sapiens	orphan G-protein coupled receptor	1570	100
		Homo sapiens	Human secreted protein encoded by gene 22 clone HFEAF41.	504	98
1296	AC004832	Homo sapiens	similar to 45 kDa secretory protein; similar to CAA10644.1 (PID:g4164418)	648	65
1297	X80035	Oryctolagus cuniculus	cysteine rich hair keratin associated protein	575	70
1298	G02645	Homo sapiens	Human secreted protein, SEQ ID NO: 6726.	223	97
1299	Y59440	Homo sapiens	Human delta3 fragment #4.	122	32
1300	W70504	Homo sapiens	Leukocyte seven times membrane-penetrating type receptor protein JEG18.	459	81
1301	Y67315	Homo sapiens	Human secreted protein BL89_13 amino acid sequence.	3916	99
1302	M77693	Homo sapiens	spermidine/spermine N1-acetyltransferase	174	96
1303	G01331	Homo sapiens	Human secreted protein, SEQ ID NO: 5412.	254	69
1304	G01491	Homo sapiens	Human secreted protein, SEQ ID NO: 5572.	747	99
1305	AF148509	Homo sapiens	alpha 1,2-mannosidase	602	98
1306	G01658	Homo sapiens	Human secreted protein, SEQ ID NO: 5739.	333	98
1307	Y90899	Homo sapiens	D1-like dopamine receptor activity modifying protein SEQ ID NO:1.	332	98
1308	AF033120	Homo sapiens	p53 regulated PA26-T2 nuclear protein	348	52
1309	Y73388	Homo sapiens	HTRM clone 3376404 protein sequence.	147	66
310	AF063243	Bos taurus	ribosomal protein L30	296	90
311	AF224494	Mus musculus	arsenite inducible RNA associated protein	688	70
1312	Y73342	Homo sapiens	HTRM clone 2709055 protein sequence.	1154	100
1313	Y99419	Homo sapiens	Human PRO1780 (UNQ842) amino acid sequence SEQ ID NO:282.	1145	78
1314	AF116667	Homo sapiens	PRO1777	433	97
1315	W75100	Homo sapiens	Human secreted protein encoded by gene 44 clone HE8CJ26.	807	97
1316	AJ272078	Homo sapiens	APOBEC-1 stimulating protein	789	100
1317	AB041533	Homo sapiens	sperm antigen	2607	98
1318	U19617	Mus musculus	Elf-1	806	92
1319	U82598	Escherichia	ferric enterobactin transport protein	768	100
1320	D90892	coli Escherichia coli	SORBITOL-6-PHOSPHATE 2- DEHYDROGENASE (EC 1.1.1.140) (GLUCITOL-6- PHOSPHATE DEHYDROGENASE) (KETOSEPHOSPHATE	709	100
1321	W67847	Homo sapiens	REDUCTASE). Human secreted protein encoded by gene 41	601	92
1200	A 127C101	IVam	clone HPBCJ74.	166	02
1322 1323	AJ276101 AJ276101	Homo sapiens	GPRC5B protein	466 504	93 97
1323	Y58628	Homo sapiens Homo sapiens	GPRC5B protein Protein regulating gene expression PRGE-21.	1584	
1324	U91561	Rattus	pyridoxinc 5'-phosphate oxidase	1277	100 89
وشاو	051301	norvegicus	pyridonine 5 -phosphate oxidase	12//	09
1326	AF125533	Homo sapiens	NADH-cytochrome b5 reductase isoform	1606	100
1327	Y32206	Homo sapiens	Human receptor molecule (REC) encoded by Incyte clone 2825826.	1531	90
1328	AF151048	Homo sapiens	HSPC214	657	85
1329	Y10530	Homo sapiens	olfactory receptor	1645	100
1330	AF180681	Homo sapiens	guanine nucleotide exchange factor	4314	99
1331	AF111856	Homo sapiens	sodium dependent phosphate transporter isoform NaPi-3b	3591	99
	l .	1			<del> </del>
1332	Y13583	Homo sapiens	G-protein coupled receptor	2171	100

SEQ	Accession	Species	Description	Smith-	%
ID	No.	}	-	Waterman	Identity
NO:				Score	
1334	Y25755	Homo sapiens	Human secreted protein encoded from gene 45.	1380	96
1335	AF152325	Homo sapiens	protocadherin gamma A5	4742	99
1336	X74070	Homo sapiens	transcription factor BTF3	639	81
1337	AF095927	Rattus	protein phosphatase 2C	1931	95
		norvegicus			
1338	G03877	Homo sapiens	Human secreted protein, SEQ ID NO: 7958.	621	100
1339	AL008582	Homo sapiens	bK223H9.2 (ortholog of A. thaliana F23F1.8)	626	100
1340	X61615	Homo sapiens	leukemia inhibitory factor receptor	5820	99
1341	Y01519	Homo sapiens	A carcinogenesis-inhibiting protein.	7528	97
1342	AF207600	Homo sapiens	ethanolamine kinase	2372	100
1343	U54807	Rattus	GTP-binding protein	1167	97
		norvegicus		1	
1344	AC020579	Arabidopsis	putative phosphoribosylformylglycinamidine	3283	51
		thaliana	synthase; 25509-29950		
1345	Y28576	Homo sapiens	Secreted peptide clone pe503_1.	944	100
1346	W74787	Homo sapiens	Human secreted protein encoded by gene 58	1171	100
			clone HHFHN61.		
1347	M55542	Homo sapiens	guanylate binding protein isoform I	2636	87
1348	AF183428	Homo sapiens	28.4 kDa protein	1329	100
1349	U70669	Homo sapiens	Fas-ligand associated factor 3	167	24
1350	AF295530	Homo sapiens	cardiac voltage gated potassium channel	562	99
			modulatory subunit		1

## TABLE 3

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino, acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
1	1351	A	2	337	1	TPSLIHQAPTPCPAGLWG/PPNGHYHGS*PGC HWPQAPHRA***GLLPPRWLGHGLPGGPAAP WAASQWVDGVAGRLPGPAWSWHASGAAPA QPGPL*LLVPGSSGLPDPRDP
2	1352	A	27	100	366	IRNSSIRPMKERETKLSAKHMITCSASYDIRGL QIETT\YHHTPIRMAKIQKT/GHHQC**ECGAT GTLIHGWWGCKVVEPLGKTVWQIPK
3	1353	A	40	3	314	·HASAHASVVLKDNSELEQQLGATGAYRARA LELEAEVAEMRQMLQLEHPFVNGADKLRPD SMYVHLNEL*QSLVENMLLTVVDTH\RTPI*R SCNYTLALILFL
4	1354	A	74	2	292	TASALFSCPDGGSLAGFAGRRASFHLECLKR QKDRGGDISQKTVLPLHLVHHQVAHTFGQAT VTCQQARQSPG*RTNPE/ALQWVLPVSDGWH VLPLP
5	1355	A	78	114	850	ENCRVASNLPGVFFSEDTAQSGSYMRISAHPP NAGGEVSNGPKRKLTLMLNFSLPSSGLNAGA FYALSTLLNRMVIWHYPGEEVNAGRIGLTIVI AGMLGAVISGIWLDRSKTYKETTLVVYIMDT GGAWWCYTFYLGTGDTCG*CFITAG\TMGFF MTGYLPLGFEFAVEL\SYPESEGISSGLLNISA QVFGIIFTISQGQIIDNYGTKPGNIFLCVFLTLG AALTAFIKADLRRQKANKETLEN
6	1356	A	81	97	376	EWFSYMLGSNMSVYHSP*SLEPLCKVLSES*A YLRVPFIRILLNAR*IRKAYKRMSLEIKLLI/RE *CLFQEMGLSLQWLYSARGDFFRATSRL
7	1357	A	93	2	872	TLSSACLIGDAWKELTIVAGAVSNQLLVWYP ATALADNKPVAPDRRISGHVGIIFSMSYLESK GLLATASEDRSVRIWKGGDLRVPGGRVQNIG HCFGHSARVWQVKLLENYLISAGEDCVCLV

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8	1358	Ā	106	3	350	VP**ARYTQGCDSGWLLATAGSD*YRGPVSL *RRGQVLGAAARG*TFPVLLPAGGSSWSRGL RIVCYGQWGRSCQGCPHQHSNCCCGPDPVS WEGAQLELGPAWL FSSLLSGRISTLRDETGAILIDGDPAACAPIIKF
						LLTEELHLRGVSIYVLRHEAQIYGITPL\VCAL LI/CRRL*SDSCMRAALNDRGLYQVLILDGLV QCLGFVDSDSRKMVSTLT
9	1359	A	115	49	186	QAWAIFKGKYKEGDTGGPAVWKTRLRCALN KSSEFNEGPERERMDV
10	1360	A	123	2	1249	KGCRTQEKVDRTEVIRTCINPVYSKLFTVDFY FEEVQRLRFEVHDISSNHNGLKEADFLGGME CTLGQIVSQRKLSKSLLKHGNTAGKSSITVIA EELSGNDDYVELAFNARKLDDKDFFSKSDPF LEIFRMNDDATQQLVHRTEVVMNNLSPAWK SFKVSVNSLCSGDPDRRLKCIVWDWDSNGK HDFIGEFTSTFKEMRGAMEGKQVQWECINPK YKAKKKNYKNSGTVILNCKIHKMHSFLDYI MGGCQIQFTVAIDFTASNGDPRNSCSLHYHP YQPNEYLKALVAVGEICQDYDSDKMFPAFGF GARIPPEYTDSHDFAINFNEDNPECAGIQGVV EAYQSCFPKAPTFTGPTNICPHSSRKVAKFRR SEGN*HQGRAFAIIFILVDPGQVGVYSQDMGP DNPGGHFV
11	1361	A	147	614	9	ACARKQLLGRTVFIWFVGQLLGGELKGYSKT NTTSSRPASSRG\TLSSSSSSSSSLTKDALPSSL KSDSTTITSGLVFPFRSLCVNPAKSSVSESVSSI KILLSSSVKYLE*KRTSCCFPDSSESKLSQLSS DERVSMGTSSRKPTNSSSSLGALKMSATS\*G SGSESPTPFFLTGLQSPPSTRPREPGLTTARNS TTLTRDC
12	1362	A	177	12	416	LIPSEPALDSLVDPRVRSRKQPFVIYPVYDTAI DTKIHFSLLDGNVGEPDMSAGFCPNHKAAM VLFLDRVYGIEVQDFLLHLLEGGFLPDLRAA ASLDT/AEIGAMDFLLS*LFTLCLMMFFFIYPFI NLLTMNVY
13	1363	A	249	535	105	WTFHRHLSPAPLIVCDQGTCVVSYYPQNIVQ MPDTQMEQGLN/HLFLDGNA*PHSVECYCPS TFEIAIKITSFVLYFHRYRAPEVLLRSSVYSSPI DVWAVGSIMAELYMLRPLFPGTSEVDEIFKIC QVLGTPKKVSTLVPKLL
14	1364	A	254	572	201	YLLTXIGNLMMLLVINADSCLRTXM*FFLGH FFFLDICYSSVTAQDAAEFPVS*KPILVWGYIT *SFFFIFSWGTNGCLLSAITYACYAAICHPLLS TMVMNRPLCTATVNATNKMGFLNSQVN
15	1365	A	257	425	68	THAKFLNKKFNIPKLVILPKLVYIVKAIPTKM AIEFLLECDQNIT\KLICENT*KNIAKNI*KRRV TFTPIET*HPVKQMIKWQ*LTAWLRNRGYKKI KQTPNSETAPSVCRNLVFDKCG
16	1366	A	263	104	481	FCIFRTTEEDRGGDDCVVSVWTKQRNNSCVK SKDVFSKPVNIFWALEESVLGVKARQPKPFFA AGNTFEMTCKVSSKNIKSPRYSVLIMAEKPV GDLSSPNETKYIISLDQDSVVKLENWTDASRV
17	1367	A	298	68	208	RKRTNNPIKLDKKFEHFKNEDI*ITSKHTKMW VSSLAMKEMLTKTTM
18	1368	A	300	904	1	LVVGITGTRHHARVIFIFLVETGFPHVGQAGL ELLTSGDPPALASQSAGITGMSHCARPKGHFG

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19	1369	A	302	3	445	TEPHSVVQAGGQWRNLSSLQAPPPGLMPLSR ISLMSSWDYRRPPQ NSPSRWAKIQMFEHTFCG*GCG/ER/NVHIHCS WICRLRPLLWRAVREYLSKLKNAELSFDPGV SLLRIYAIDMPTSI*DEKEALLFAFLAFHE*HC KSRIWAVIQ/CIHLWDWLRKL*CFHRMKFYA
20	1370	A	304		1339	AV*NKPRHLLSHIWKDVQNILLK  FFFCGKEVPLFEQNKHPGPRATTSPGA/HARA LLSAGEFTAGVGLSP*AIHSFVWLCTFIQHGA GGPCHQPGGSPGPWMHTIQAGHLWEGAYPG GSSTWHQVPGQLGGSWGPRERSLLGSFIKCSP CPHPPGFRLWMSPNQKPTENPGVMGRVWR LMPGESPLIWEAEGKEDHLSPEGQGIISE/PVA PLHSSLGNTVKP*PKNQKPKQNRSRHGQ\GF MAGQGQSRPAAR*PPCPALTPASHSAGTWPP RICRTVPGGPCPSPSGFRSCRR*GFSA*TRSWP DAEPPSTPDTAPRCCTQSDTSSQGPQ*S*WRR CRALPGRLCSAPAAGLRRARPRLSESRRGNSP PASPAAASARCPSWGPSCPARPPSRPAAGTEP AAPSRCTAWLRGEREPGPRPPGRRPRSGRGP VSFAPEVLSLPAVRQTKSWRWRNEEEITRPW ALVRSRGG
21	1371	A	326	799	1587	GSQVLPPPPSQDSATLPQDA*GPRAAPGQPVC E*GLQGAGVRRLRGEVLCQPQP*GAL*EQCLP HLSFSPRQGAAPDTEPSAWGPAPTGATGPGLP LRHVRLFSAGAPRGAATFCPPALLHGPAWPP ARPMFRGHPPVRPLGPWGKVAAGPRALCLA GVPAVQGECATKPSG*GL*PAHLRGPPGPEVL QWHWQLSAGRDPVPAEDPPL*EGPLGPGGPA AAQAEPGADPEPEDKDQAAESRPAGAMSLSA QGSGPVGGQGLR
22	1372	A	327	146	652	PHLENPHPEHSFPGAPLT*STLSWSILSPREPSP GAPCYPGHPHLENPHLEHLLTWRTVTWSTLL PGAPCYPEHPHLEHPLTWSTPHLEHPSPGEPL SCRTPTRSILHRDHPLP*CLSTEESPI*GWGSLP APPSTPLVLDVAPPGPQPASSCPGRDSCYSVP GTVVSP
23	1373	A	348	397		CIVSSCQGTRKPCHLEDANKINKQSPTLEKIES LQESL*VKQ*LIVAEKYVQILHPRKKYFQRPL NNEKRKMKKRKEEKKKCRERMQRRSKWRR EEKKE*RREE\EERKKEKEDRKERRKETSPRG SRRLLRD
24	1374	A	362	170	352	GRALDTAAGSPVQTAHGLPSDALAPLDDSMP WEGRTTAQWSLHRKRHLARTLLVSRVRGPQ
25	1375	A	384	373	128	YLITTILETGYLWKNRHSDQ*KRTENPERDQH KYPKVDFCKSNSMKNRLCNKWHWTNWIFTD KKINLNLKPHTKLTPNIKKN
26	1376	A	397	383	165	EVKNTNPFIFSGTNLTIWIRSI*RKSDEINQRTK *MEKYSISLDRRLNTVKMSFLPNLIYKFNTISI KIPANF
27	1377	A	406	103	380	KSKATGYMVNI*KLIV\FLYANDEQLEIEMNK IVP\FNGSKNKIAFTNLTKYQNIQNRHAENYKI LVNKIEDLNKWRNVLLSWIGRRNIINTMT

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28	1378	A	408	sequence 14	427	nucleotide insertion TICTNKFNNLDEIK/FLERHKLSKLTQEEVENL ITLKTSRETELVINK*VIPHKEKPGPDSFTGEF YQTFKEEL/IJ/ILHKLFQTIKYGRILPNSVYETSI TLKPKPEKDL\KENYRPLPLSNIDAK\LNKTLA NRI**HIR
29	1379	A	434	395	128	IYSKMCMERQRLNN*ILKKNKVRGIAVPDVK VYYKPTVIK/TSWIL*KDSHIVEWNRLENLEID PN/IKRLILDKGAEATEWRKDSFFRQWQ
30	1380	A	455	2	228	FFFETESHSVTQAGVQWCNPGFKRFSCFGLSS SWDYRYAPPRP\ANF\*FLVETGFYYVAQAGL KLLSPGDLPALAS
31	1381	A	462	393	2	QLMFDKGVKNIH\WGWTPPFTK*YWKNWISI CRRMNLNPYLSRYIKINSR\KDLTVRPEPIKLV EENTGKTIQDTGLGK*FIAKTSKAQSTKTNK* KRQTRYIKLK\KKSTASKENNRVKRQPLE*EK IFAN
32	1382	A	474	125	471	VKPYEIAVFLVKPIEYK*HLLSDPAIPLSGI*LK EIKAYT/RRICTPMFAAPVSVIA/RN*KQSK/CQ KQ*YVHRMEYYTTIKRSEILICTTTWVDFRNT ILRETDRIHKTTYDVISLI
33	1383	A	488	1825	2	KSACSFICSEEQPASPSPLKPGTYASET\RPRDP HAAGPRRDSSEAETRRPRGA/DGSGTVVKGT PGSPAPPCSWGHGG\ETEGAG*CPAAPGTDLR APGGSAGS*\GLPSAGGSRGRKGWRAAGRQP STR*GRPGRHGGRGE*AGHPEPRQSALQSAG L/ASSPEPMGAALAEDGSGDSRGAGPRPQE*P PSVLSRS\GS*G*G*AASGTASSPRSHSSRLGPP SAGFHGLRCGQPPFAAAPPGPWPGTGRPAGG AGSPPAAAGTAPPATRGAQSRRQNRTAGRNA SPQTAAGAGSPVQWALSRATG*TGETGSWC AGGTHQATHLTAAWVCPPTWSVRPGGSGPA AGLGR*GRHPAQSPPLPVPRG*PAWPQEAPSP SPASSEVALSSGSCWPDQAPGPARGSPPAPLA PAWPAAGRGRQR*GRQSAHPPPRR*STAVSL SGTS*WRRSP*AGTRTQQC*SPWLVPACSSRP L*RGTRRPSTQQSPQTTGTPGRSAGPGHPRS* GGRSPAGTGHLGAQTVASPH*GHWPTALSCL WASASPPGPEAPPQTGACIGTNCRYRAASAR RSSVAPACA*GWQ*AGSPPAVLRGPP*RVRER GALTHRPRAPDE
34	1384	A	497	422	2	APGASVGRAQAAEG*RGGPTGRPPSALGVS/E AGRAGRAGEGRPVPPAYPLCKSAQTSGPPKA RLS\PPLASCGGRGPPGGAACATCAPPAGPAR SSRCRRRSPPE*GPR*PSRPARPSPGSAASRRQ KLTPCRCQFRGLCA
35	1385	A	509	156	475	PTPYPGE*QAAFLLRGPGLRPPA/DPSLR/HRN LTELVVAVTDENIVGLFAALLAERRVLLTAS KLSTLTSCDHAFCALLYPMRWEHVLIPTLPPH LLDYC*CPPLPRT
36	1386	A	. 512	3	1631	FFFSFVCHLYCVSPTPGPHGRLATWL/PGLLA FLGLAAGGQTLCPAGELPGHARAQASGAPGS VLIAVPGRRRVHTCGPGPAAPSTRGECPPPAL, GHTRPARPRPV\PFAPAVPQEPGGQGHGAA/P PATGHSAPRGCPPARAAPTGSATPAPPPAACA AFHSAWSVPPAGRQQG*RVPAPAFRRTTPGT PGQHLLDRPGAPPAQGSGPAPAPPPRLAGPA GPAAPPPGPPAASWHSSLSKSSSSL\GWSPPLP VGPGSLQ*TPPPQGPHLSGSCGGTSSWRGQR AAVARRLRSWNACGLSRVAGRSSASYPGRE

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37	1387	A	620	828	1	FRLPLAAGA/RGAAEPRVAVSMAPDPSAKIH WEASPEMQSKCHQKGKNNQTECFNHVRFLQ RLNSTHLYACGTHAFQPLCAAIDAEAFTLPTS FEEGKEKCPYDPARGFTGLIIDGGLYTATRYE FRSIPDIRRSRHPHSLRTEETPMHWLNG*EDE AQDDGG*GTISSFLLPWPADHPTPKSPGEPVH SIPVCCQVRGQPQSGGKESPACLKSLSNCLTH \DAEFVFSVLVRESKASAVGDDDKVYYFFTE RATEKESGSFTQSRSSHRVARGIPPL
38	1388	A	739	1	427	FRAMVSSTLKLĞISILNGGNAEVQ/QGNRĞKĞ TSEEGKEG*EVPV*LPVSPPLPRPLQKMLDYL KDKKEVGFFQSIQALMQTC\GEKVMADDEFT QDLFRFLQLLCEGHNNDFQNYLRTQTGNTTT INIICTVDYLLRLQESI
39	1389	A	767	I	1030	TLDLTGPLLLGGVPNVPKDFRGRNRQFGGCM RNLSVDGKNVDMAGFIANNGTREGCAARRN FCDGRRRQNGGTCVNRWNMYLCECPLRFGG KNCEQGEWPASSIPPVTAAWEALLLDVPGTT VRGLHIQVRQPLVVYAAFTVDSHRPLQETVL RRAPAPASGVPSPSGVGWDR*AGPAEPSPSTP ATVIISVPWYLGLMFRTRKEDSVLMEATSGG PTSFRLQVTGAPCHQGTC*VGARGRDPMLSG LRVTDGEWHHLLELKNVKEDSEMKHLVTM TLDYGMDQVSWHLHLLWG*TLPPAQGKTGA SEDKVSVRRGFRGCMQVRGGCGGRGEACPS OAAPRL
40	1390	A	801	69	399	IHKIIIHKEDLNKWKYILCSGMERLSTVMIPVV PQIIYKFNA*Q\VILKFTW*E*GAKITILRKNKL RGLVLVPLSTC*VKYLLDKVLPHIKTYYEAR VNKSVVLVQVTIM
41	1391	A	835	7	195	SMLKERKVFQFPSCLFFQYITWLGPPYHVLFD SSVTNFSIGAK*DILQSVMNCLYAKRIPCVT
42	1392	A	841	1	415	GSTHASGYDKTPDFILQVPVAVEGHIIHWIES KASFGDECSHHAYLHDQFWSYWNSLKHRTW QGIGTVASNLSQL*TLNAPFPELLLFRSLARTG FVLT*\RFGPGLVIYWYGFIQELDCNRERGILL KACFPTNIVTL
43	1393	A	845	358	92	PALSPAPVPQKKGSPLPLDPCLGPSSWLLSVG LGWPRL*PRRGPGDPGSLPATPPLLTPPHTLLP QRPMLPPSHAGLARPPPPEPISVP
44	1394	A	853	452	1	LPQYCFFPRLSPKSKLVKHSAL**PSALKPPTK SPRCIPRTSLYFTICC/PPALQL/SPIEDPPAIYRS PPTHMLRSASQPLNQAPTLVKGHPPSRFLQG QVSCPPQPTLPREKPLPLHLRPPPRPAQPPLPR PLTFSTRRNVDPEIPERFR
45	1395	A	894	379	162	GVYPPTVFDNYSVQTSVDGQIVSLNTWDTAG QEEYD/RLRTLS*PQTSIFVICFSIGNLEFPIYGT WLSMSMGK
46	1396	A	900	1	366	TTKKTLISNNVSSRSLPILPELKAFSLAFNDPL EIQKYMRT/DQ*CVTHDISLYIVTKLALIFLIPR VFLFHQLNIT**CLHFFTMTTFIAIPFSFLFLGR

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47	1397	A	944	162	2	QLQNLASRGCL*SQLLRRLRRENRLNPGGGG CSEIAP\CTPAWVTQRDFFRKKK
48	1398	A	963	216	308	HFTPDRIAIVKNTRDSHCWRGC*EEGAPARC
49	1399	A	967	466	1	PRKRESWWGERLP/PRGFPPAAEDAPAPGWK GRKHASRTARAHVFHPIRQSIRSPVRGRPGDP RAAHTRSAGTRLQCKASRGG*GKGPAPTR*E GGPGSAPAPLPASSGCSLFPDSSPWTPPPPAPG AAAAQP**TPRCPAALRAGAHIGRVGRPY
50	1400	A	973	45	421	EKCIQALDVFVFCYIDHSSHCLMSCD*E/DQA LNFMPLEMEPKMSKLAFGCQRSSTSDDDSGC ALEEYAWVPPGLRPEQIQLYFACLPEEKVPY VNSPGEKHRIKQLLYQLPPHDNEVRYCQSLSE E
51	1401	A	992	2095	194	IRIRHEAARSCLGCAAGHVPAPGLRLLPTVRG PPGRRGPAAPGCVCY*SGESTFVSHVPQRMA WPGSAPPRGFHPLQSQTSPSDTVSSPQLSKEE DGPGWEHPLSSSL*SLGQAGGNH*QPEELAG WEPRGPPSLAPSSPT/TMWTALVLIWIFSLSLS ESHAASNDPRNFVPNKMWKGLVKRNASVET VDNKTSEDVTMAAASPVTLTKGTSAAHLNS MEVTTEDTSRTDVSEPATSGVAADGVTSIAPT AVASSTTAASITTAASSMTVASSAPTTAASST TVASIAPTTAASSMTAASSTPMTLALPAPTST STGRTPSTTATGHPSLSTALAQVPKSSALPRT ATLATLATRAQTVATTANTSSPMSTRPSPSKH MPSDTAASPVPPMRPQAQGPISQVSVDQPVV NTTNKSTPMPSNTTPEPAPTPTVVTTTKAQAR EPTASPVPVPHTSPIPEMAMSPTTQPSPMPYT QRAAGPGTSQAPEQVETEATPGTDSTGPTPRS SGGTKMPATDSCQPSTQGQYMV/DHH*APHP GRGRQNSPSGGAVTRGDPFHHSLGFVCPAGL *ELQEEGLHPGGLLNQRDVCGLRNVRGAGA WREAWPLPRPFLLPLRPNQVLPNSFGAIEEIC QMLKHI
52	1402	A	994	1	462	ESGEFLVSFTLKKPTNVFHHINGMKFFNK/LIF *SHTDIAFYKIQHPFMLKALTKWA*EGT*PDR RYLH*SLRLNGEQLKTFPLRSGMR*G/CAILPL VLNAMLSIVPAVVPAGKTRHEKEITCPLIGQE EK*FS*FVGDMNTCVENKKESKKLLE
53	1403	A	1011	1	630	PEVIQQSAYDSKADIWSLGITAIELAKGEPPNS DMHPMRVLFLIPKNNPPTHCWRRŁLESFKEV *LMLA*TKDPSI\RPTAKELLKHKFIVKNSKKT SYLTELIDRFKRWKAEGHSDDESDSEGSDSES TSRENNTHPEWSFTTVRKKPDPKKVQNGAEQ DLVQTLSCLSMIITPAFAELKQQDENNASRNQ AIEELEKSIAVAEAAGPG
54	1404	A	1016	1	222	ISIDA*KAFDKIQH/CFMITTLKKLGIDGKYLN TIKAIDDRHTVSTILNVEKLKAFL*RSGTRQRF PISGSGARI
55	1405	A	1033	3	366	HASVDGDEGSDDVYYYYTPAILRELQALNTA EAAEHRPEEDRMLSEDPWRPAHMIKGYMPL HNIPHTEVIDVTGLNQSHLYQHLNKGTPMKT QKRAA\LYTWHVLEQLEILRQINQQSHGPG
56	1406	A	1044	5	429	SVLTLQTRSPSKPLS\RKLMDWEVVSRNSISE DRLETQSRASRSPPVTPNQSQETPVDGKPLAL PPNQSQKNIRYHIHYLHLQYYLDRHISATLPIP SSSGIPTPIAVITDALTDLVELILGQPCSEESGR APGTLFLLAL

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57	1407	A	1050	11	430	GAYAFETNGFPIMLVLTTDKIEGDVGIAGLYD MH\ISLPMAFLLRTLVRCTSYIIPVTHVLSTPV TCLRRREKDGVIVDVLSDTASNHNGFPVEEH ADDTHPARLQGPTLRSQPMGPLKHKAFEERA NLGLVQRRLRLED
58	1408	A	1058	258	419	LKHRDTPVVGANNRALSCTPLTSLTLCALCPL PCLGCPTXATCRLYQTTVAVVF
59	1409	A	1064	3	425	KAFSFTTSLIGHQRMHTGERPYKCKECGKTF KGSSSLNNHQRIHTGEKPYKCNECGRAFSQC SSLIQHHRIHTGEKPYECTQCGKAFTSISRLSR HHRIHTGEKPFHCNECGKVFSYHSALIIHQRIH TGEKPYACKDVGK
60	1410	A	1065	204	419	GGPPGPFLAHTHAGLQAPGPLLAPAGDEGDL LLLAVQQSCLADHLLTASWGGK/DPIPTKALG EGQEGLPLTV
61	1411	A	1079	3	383	RHSRAHLCQPFHLVMRDLLQLGQDIPQGCHY LEENHLIHRDIAARNCLLSCAAPTRAATIGDF GMARYIYRTRYYQLGDRAL/LPRKWMPPEAL LEGIFTYNTDSWTFGVLLWEIFSLGYMPYPGR TN
62	1412	A	1080	1	859	VVEFLWSRRPSGSSDPRPRRPASKCQMMEER ANLMHMMKLSIKVLLQSALSLGRSLDADHA PLQQFFVVMEHCLKHGLKVKKSFIGQNKSFF GPLELVEKLCPEASDIATSVRNLPELKTAVGR GRAWLYLALMQKKLADYLKVLIDNKHLLSE FYEPEALMMEEEGMVIVGLLVGLNVLDANL\ CLKGEDLDSQVGVIDFSLYLKDVQDLDGGKE HERITDVLDQKNYVEELNRHLSCTVGDLQTK IDGLEKTNSKLQERVSAATDRICSLQEEQQQL REQNELIR
63	1413	A	1083	2	615	SSFAKHKRIHTGEKPFICLECGKAFTSSTTLTK HRRIHTGEKPYTCEECGKAFRQSAILYVHRRI HTGEKPYTCGECGKTFRQSANLYAHKKIHTG EKPYTCGDCGKTFRQSANLYAHKKIHTG\EKP YKCKECGKAFKSYYSILKHKRTHTRGMSYEG DEC/QRSLN/RSSILSNHKIIHNEEK/PLKCEKCE KAFNHTSICCRHKKN
64	1414	A	1084	946	1	KKQDLSSSLTDDSKNAQAPLALTESHLATLA SSSQSPEAIKQLLDSGLPSLLVRSLASFCFSHIS SSESIAQSIDISQDKLRRHHVPQQCNKMPITAD LVAPILRFLTEVGNSHIMKDWLGGSEVNPLW TALLFLLCHSGSTSGS\HNLG\AQQDQCKISFS FFSWLTTGLTTQQRTAIE\NATVAFF\LQCI\SC HPNNQKLMAQVLCELFQTSPQRGNLPTSGNI S\GFIR\RLFLQLMLEDEKVTMFLQSPCPLYKG RINATSHVIQHP\MYGAGHKFRTLHLPVSTTL SDVLDRVSDTPSITAKLISKQKDDKKKK
65	1415	A	1087	103	324	PRAFEFVHTEMIVG/RVQNIHLFTLQVLEDRA LFTMSVGSSLWSTYLIHVMALP/DRELLKPNA SVALHKLSNALV
66	1416	A	1095	3	493	HETCSVTHIVSFSLPFLNPSHPASTPGHTENEQ PSLVWFDRGKFYLTFEGSSRGPSPLTMGAQD TLPVAAAFTETVNAYFKGADPSKCIVKITGE MVLSFPAGITRHFANNPSPAALTFRVINFSRLE HVLPNPQLLCCDNTQNDANTK\EFWVNMPNL MTHLK
67	1417	A	1098	57	356	LKLTSLGFIIGVSVVGNLLISILLVKDKTLHRA PYYFLLDLCCSDILRSAICFPFVFNSVKNGST WTYGTLTCKVIAFLGVLSCFHTAFMLFCISVT

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
68	1418	A	1106		1326	MGKISATGINMGTKCSWALVWHLESYDPKH YEREGMQDWKTASGQSEEATQQSSQKPQPH YTTYQSSSFLKYSSESHLLAWRENSSEGSFQF PGRSRARPPRTRQQRRGAAAGPGRGAVRLG HPQSAAQPQLRAAARIPESPAAFPAQPRPGSA RNSDASGPASLSRTLGRASSPRPPQAPDVTAP SPAALAPRAARGGSRAAALAGAEAEEPLRTL APRPTRAAAPPPPPPPPPPPPPPPPPPPVRCVSR RARAPPWR/PAATGPPPNRPVAPSRKLGSARAP APALQIRKGTSSGLPGRGGGSGPGNNLSSVA GNWRGSSFAVERPGMAKYQGEVQSLKLDDD SVIEGVSDQVLVAVVVSFALIATLVYALFRNV HQNIHPENQELVRVLREQLQTEQDAPAATRQ QFYTDMYCPICLHQASFPVETNCGHLFCGSLT PNSIW
69	1419	A	1107	2	466	FDTARLHEFGTSITQIFAVDNREDLQKWMEA FWQHFFDLSQWKHCCEELMKIEIMSPRKPPLF LTKEATSVYIIDMSIDSPMKLESLTDIIQKKIEE TNGQFLIGQREESLP/SS/CGPHSLMVTIKWSS RKRY/SYPASEPLHDEKGKKRQAPLPPSDK
70	1420	A	1111	698	23	ALRRLHYVRATKV\FLSFRRPFWREEHIEGGH SNTDRPSRMIFYPPPREGALLLASYTWSDAAA AFAGLSREEALRLALDDVAALHGPVVRQLW DGTGVVKRWAEDQHSQGGFVVQPPALWQT EKDDWTVPYGRIYFAGEHTAYPHGWVETAV KSALRAAIKINSRKGPASDTASPEGHASDMEG QGHVHGVASSPSHDLAKEEGSHPPVQGQLSL QNTTHTRTSH
71	1421	A	1119	2	385	QKQTLQNGYLDSSMDILYLGSLPPELQVSSDE PPGPPEQAGLSQFHLEPETQNPETTEEIQSSLQ QEAAAQLPQLPEVVELSSTKA\EAPALPSQSL EGVHSSTEQKAPAQQLPAFEEILAPLLIHHE
72		A	1127	1	906	HAQYVGPYRLEKTLGKGQTGLVKLGVHCIT GQKVAIKIVNREKLSESVLMKVEREIAIL\RLI EHPHVLKLHGVYENKKYFPPDELTSGPSMLA QVSPHGKLSARRSWDLLSGFPRYLVLEHVSG GELFDYLVKKGRLTPKEARKFFRQIVSALDFC HSYSICHRDLKPENLLLDEKNNIRIADFGMAS LQVGDSLLETSCGSPHYACPEVIKGEKYDGR RADMWSCGVILFALLVGALPFDDDNLRQLLE KVKRGVFHMPHFIPPDCQSLLRGMIEVEPEKR LSLEQIQKHPWYLGGNFIS
73	1423	A	1128	l	802	LRNALDVLHREVPRVLVNLVDFLNPTIMRQV FLGNPDKCPVQQA/MLEPLGSKTETLDLRAE MPITCPTQNEPFLRTPRNSNYTYPIKPAIENWG SDFLCTEWKASNSVPTSVHQLRPADIKVVAA LGDSLTTAVGARPNNSSDLPTSWRGLSWSIG GDGNLETHTTLPNILKKFNPYLLGFSTSTWEG TAGLNVAAEGARARDMPAQAWDLVERMKN SPDINLEKDWKLVTLFIGGNDLCHYCENPEA HLATEYVQHIQQALDILSE
74	1424	A	1139	60	480	FREPCLLVPGDHQPLREASWLA/LPPIGLWGT DSPLCCVEVAIPCNKGAHSVGLKGWLLAQG VLGMRDTIPQEHPWESTPDLCFCRDPEEIEVE EQPAADAAVAKGEF/QGEQIAPVPA\IIAAHPE AADPAPVHTTAHPKGA
75	1425	A	1147	2	413	PFPHQHPQEP\KGSCWPQSALRGQCPGPVLGV TTTSDLCSLQVPVSSHRNPLLDLAAYDQEGR

SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence	1	09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence		]	914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
	1	l	i	amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
1	l	ł	ł	residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
		l		peptide		/=possible nucleotide deletion, \=possible
	L			sequence		nucleotide insertion
	1	1	İ			RFDNFSSLSIQWESTRPVLASIEPELPMQLVSQ
			1		ļ	DDESGQKKLHGLQAILVHEASGTTAITATAT
L		l	i	ĺ	ì	GYQESHLSSAR
76	1426	A	1155	38	410	PIISAPAQDDPILLSFIHCLHANLLCVWRRDVK
			1			PDCKEIWIFWWGDEPNLV\VQYIMNCMLWK
l	1	Ì	l		}	KDSGKMAFPMNVGRC/FFKEIHNLLERCLMD
			1			KNFVLIGKWFVRPYYKDEKPVNKSEHLSCAF
						T
77	1427	A	1162	526	350	RFPQGLEDVSTYPVLIEELLSRGWSEEELQGV
						LRGNLLRVFRQVEKVQEENKWQSPLED
78	1428	A	1171	I	1293	MAESASPPSSSAAAPAAEPGVTTEQPGPRSPP
1.0	1 .20	^^	1	_	1275	SSPPGLEEPLDGADPHVPHPDLAPIAFFCLROT
		Ì				TSPRNWCIKMVCNPWFECVSMLVILLNCVTL
						GMYQPCDDMDCLSDRCKILQVFDDFIFIFFA
[	[	1	[			MEMVLKMVALGIFGKKCYLGDTWNRLDFFI
			1			VMAGMVEYSLDLQNINLSAIRTVRVLRPLKA
1	ł		l			INRVPSMRILVNLLLDTLPMLGNVLLLCFFVF
			l		İ	
						FIFGIIGVQLWAGLLRNRCFLEENFTIQGDVAL
(			1			PP\YYQPEEDDEMPFICSLSGDNGIMGCHEIPP
1	ĺ	ĺ	1			LKEQGRECCLSKDDVYDFGAERQDLNASGL
			ļ			CVNWNRYYNVCRTGSANPHKGAINFDNIGY
1	Ì		i			AWIVIFQVITLEGWVEIMYYVMDAHSFYNFI
	ŀ	ļ			1	YFILLIIVSVREPGLLGGSFSTAQSPKCQGDSFP
		ļ				GVAAESLLLRGWVLWLPGGG
79	1429	A	1175	1	405	PNDFFKDMFPDLPGGPLGPIKAENDYGAYLN
	·	[	[			FLSATHLGGLFPPWPLVEERKLKPKASQQCPI
						CHKVIMGAGKLPRHMRTHTGEKPYMCTICE
1	!	ļ	<b>!</b> ,			VRFTRQDKLKIHMRKHTGERPYLCIHCNAKF
						VHNYDLKNHMR
80	1430	A	1182	25	198	EMNELSQQLSQQGGRGASQCPSPPAPTLPNPT
	!		[			PLCQLQLQRVNTGLPTPPCHPGAGAA
81	1431	A	1186	254	583	KTVLDVGAGTGILSIFCAQAGARRVYAVEAS
		}	1			AIWQQAREVVRFNGLEDRVHVLPGPVETVEL
1		1	ł i			PEQVDAIVSEWMGYGLLHESMLSSVLHARTK
						VVKDGGFFLPXSSELFM
82	1432	Α	1187	2	716	DFVDAARNLPLESTKSPAEPSKSVPSLE\DPRA
						SSQGLPSQGPVQNQGRRGEQRPKKF/TVIQHT
						SSFEKSDSLEQPSGLEGEDKPLAQFPSPPPAPH
1					{	GRSAHSLQPKLVRQPNIQVPEILVTEEPDRPD
						TEPEPPPKEPEKTEEFQWPQGSQTLAQFPVEK
			]			LPPKKKRLGLAKMAQSSGESSFESSVPLFRSP
						SQESNVSLSGSSRSALFERDDHGKAEAPSPSF
					-	DMGPKPLGTHMLTV
83	1433	A	1188	517	804	ESPGLSKVLRTGAFAYPFLFDNLPLFYRLGLC
ری	1433	, A.	1100	317	004	,
						WGRGHGCGQEALSTSHGYHLFCALLTGFLFA
1						SHLPERLAPGRFDYIGHSHQLFHICAVLGTHF
	1.40:		110-			Q
84	1434	A	1192	45	476	LGDVGFWVERTPVHEAAQRGESLQLQQLIES
						GACVNQVTVDSITPLHAASLQGQARCVQLLL
}				İ		AAGAQVDARNIDGSTPLCECLRLGQHRVCEA
1						LAVLRGQGQPSPVHSVPPARGLHXREFRMC*
						GFLFDVGXNLEAHEFHFGEP
85	1435	A	1194	69	410	KRSEEASAPPFPLGGTGAAPTRASLPEQILLPR
						SCLEARKSQPDEKLLSALHNSRTWN*EPRRSQ
}						HRLVSPEVHPGRRGSSPGVAECKLTSAYFRT
}						GRSPCPSLPGTTRTNSLL
86	1436	A	1215	3	405	LPSHTCGNPGRLPNGIQQGSTFNLGDKVRYSC
				-		NLGFFLEGHAVLTCHAGSENSATWDFPLPSC
1 1		'				RADDACGGTLRG/AEWHHLQPPLPLG/ATKN
					<u> </u>	THE DATE OF THE WITH VELLE TO ALKIN

SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide	)	in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-	ļ	USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence	İ	09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence	[	[	914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
		Ì		amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
		l		residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
	ł	İ	1	peptide		/=possible nucleotide deletion, \=possible
<u> </u>		ļ	<del> </del>	sequence		nucleotide insertion
		Ì				NADCTWTILAELGDTIALVFIDFQLEDGYDFL EVTGTEGSSLW
87	1437	A	1216	226	964	GTARFGPMVGFGANRRAGRLPSLVLGVLLV
] "	1,	1 11	1210	] 220	507	VIVVLAFNYWSISSRHVLLQEEVAELQGQVQ
]	ļ	]	j	ļ	j	RTEVARGRLEKRNSDLFAVVGHAQETDRPEG
	i '	1	ļ			GRLRPPQQPAAGQRGPREEM\EDDKVKLQNN
		ŀ			Ì	ISYQMADIHHLKEQLAELRQEFLRQEDQLQD
ľ	i ·	Ì	(		ĺ	YRKNNTYLVKRLEYESFQCGQQMKELRAQH
	ļ					EENIKKLADQFLEEQKQETQKIQSNDGKELDI
1	}		1	}	l	NNQVVPKNIPKVAENVADKNEEPSSNHIPHG
88	1438	A	1218	1	534	PEFGTTISCGYLMATDVSRRPSVHKAVEIEQE
						RVKSAGAWIIHPYSDFRFYWDLIMLLLMVGN
1	J .	]	]		}	LIVLPVGITFFKEENSP\PWIVFNVLSDTFFLLD
	1		l			LVLNFRTGIVVEEGAEILLAPRAIRTRYLRTW
			1			FLVDLISSIPVDYIFLVVELEPRLDAEVYKTAR
			<u> </u>			ALRIVRFTKILSLLRL
89	1439	A	1223	1	743	MGFDEVFMINLRRRQDRRERMLRALQAQEIE
						CRLVEAVDGKVGMLTRSNAAPGRHLAMLET
	) S-	)	1		}	LVVVAPRFVDADNLILNPDTLSLLIAENKTVV
İ						APMLDSRAAYSNFWCGMTSQGYYKRTPAYI
			1			PIRKRDRRGCFAVPMVHSTFLIDLRKAASRNL
	ſ	ſ	j .		<b>[</b>	\AFYPPHPDYTWSFDDIIVFAFSCKQ\AEVQMY
1						VCNKEEYGFLPVPLRAHSTLQDEAESFMHVQ
90	1440	A	1227	2	349	LEVMVPSSPSSAQSMAVVSADHIGLVISYL
90	1440	A	1227	2	349	NKTSFIFYLKNIVVADLIMTLTFPFRIVHDAGF
			1			GPWDFKFILCRYTSVLFYANMDTSIVVLGLIT/ YDRY/WKVVRHL/WDSWMTGI/SFTRVYLLG
1 .					}	LGARLVWFGKLILAKGGHGGISWL
91	1441	A	1245	3	1937	LGSSDVRAPQRSELGAESPSRMVASQAYNLT
	1.4.1	12	12-13	3	1557	SALTPILTRSRVLNEEPLTLAGF\SRAPANLSD
			J	,	j	VVQLIFLVDSNPFPFGYISNYTVSTKVASMAF
		Ì				QTQAGAQIPIERLASERAITVKVPNNSDWAAR
						GHRSSANSV\VQPQAFVGAVVTLDSSNPAAV
i l					}	LHLQLNYTLLDGRYLSEEPEPYLAVYLHSEPR
			Ì			PNEHNCSASRRIRPESLQGADHRPYTFFISPGT
1						RDPVGSYRLNLSSHFRWSALEVSVGLYTSLC
1						QYFSEEDVVWRTEGLLPLEETSPRQAVCLTR
						HLTAFGTSLFVPPSHIRFVFPEPTADVNYIVML
					•	TCAVCLVTYMVMAAILHKLDQLDASRGRAIP
			]			FCGQRGRFKYEILVKTGWGRGSGTTAHVGIM
						LYGVDSRSGHRHLDGDRAFHRNSLDIFQIATP
			i			HSLGSMWKIRVWHDNKGLSPAWFLQHIIVRD
			[			LQTARSTFFLVNDWLSVETEANGGLVEKEVL
						AASKASFRVPTPS\AALLRFRRLLVAELQRGF
						FDKHIWLSIWDRPPRSCFTRIQRATCCVLLICL
					}	FLGANAVWYGAVGDSAYSTGRVSRLNPLSV
						DTVAVGLVSSVVVYPVYLAILFLFRMSRSKV
]			]		]	GWGWGPGSTGNGAWASAPCPEPPLSSAAAR GKGVHQRLLGKGQHT
92	1442	A	1246	5	562	VFDEENILNELNDPLREEIVNFNCRKLVATMP
12	1774	Λ.	1240	J	502	LFANADPNFVTAMLSKLRFEVFQPGDYIIREG
					}	AVGKKMYFIQHGVAGVITKSSKEMKLTDGS
				·		YFGEICLLTKGRRTASVRADTYCRLYSLSVD
				, i		NFNEVLEEYPMMRRAFETVAIDRLDRIGKKN
			)			SILLQKFQKDLNTGVFNNQENEILKQIVKH
93	1443	A	1249	180	901	TVPPPPGGPSPAPLHPKRSPTSTGEAELKEERL
			~- '-	•••		PGRKASCSTAGSGSRGLPP\SSPMVSSAHNPN
						KAEIPERRKDSTSTPNNLPPSMMTRRNTYVCT
			[			ERPGAERPSLLPNGKENSSGTPRVPPASPSSHS

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion LAPPSGERSRLARGSTIRSTFHGGQVRDRRAG
		}				GGGGGGVQNGPPASPTLAHEAAPLPAGRPRP TTNLFTKLTSKLTRRVADEPERIGGPEVTRRP RQEDHLSPGGRGCSEL
94	1444	A	1261	3	385	KFSQWGLTKPKLSNASP/WISLVKKLMKKWS VTQNLTFREQLEAGIRYFDLRVSSKPGDADQ EIYFIHGLFGIKVWDGLMEIDSFLTQHPQEIIFL DFNHFYAMDETHHKCLVLRIQEAFGNKLCPA CR
95	1445	A	1282	2	550	GPRDNPG\EDPRFEIVEHFGIAWFTFELVARFA VAPDFLKFFKNALNLIDLMSIVPFYITLVVNL VVESTPTLANLGRVAQVLRLMRIFRILKLARH STGLRSLGATLKYSYKEVGLLLLYLSVGISIFS VVAYTIEKEEN\EGLATIPACWWWATVSMTT VGYGDVVPGTTAGKLTASACILA
96	1446	A	1294	1	1456	QLLPPSNRENAGLLVGRCLCSAALRPVGDLIT SSGQVAVRNAPQAGSAKAGKGKFQDNFEFIQ YFKKFFDANCNEKDYNPVAAGQGQETEVAP SIVAPVLNKPNQCPEGYICVKAGRNPNYGYT SFDTFSWAFLSLFRLMTQDYWENLYQLTLRA AETTYMIF/LV/LVILLGSLYLVTLILAV/VAMA YEEQNQATLEEAEQKEAEFQQMLEQLKKQQ EAAQQAATATASEHSREPSAAGRLSDSSSEAS KLSSKSAKERRNRRKKRKQKEQSGGEEKDED EFQKSESEDSIRRKGFRFSIEGNRLTYEKRYSS PHQSLLSIRGSLFSPRNSRTSLFSFRGRAKDV GSENDFADDEHSTFEDNESRRDSLFVPRRHGE RRNSNLSQTSRSSRMLAVFPANGKMHSTVDC NGVVSLVGGPSVPTSPVGQLLPEVIIDKPATD DNGTTTETEMRKRRSSSFHVSMDFLEDPSQR QRAMSIASILTNTVE
97	1447	A	1295	2	2057	IQTQLPTKSSQQLRKGGNCVRCKMQMNFIAE EVLLKYRITFYNNNKGPNMLYIEIKAFVHFMI NRYLSYGSGPKRFPLVDVLQYALEFASSKPV CTSPVDDIDASSPPSGSIPSQTLPSTTEQQGALS SELPSTSPSSVAAISSRSVIHKPFTQSRIPPDLP MHPAPRHITEEELSVLESCLHRWRTEIENDTR DLQESISRIHRTIELMYSDKSMIQVPYRLHAV LVHEGQANAGHYWAYIFDHRESRWMKYNDI AVTKSSWEELVRDSFGGYRNASAYCLMYIN DKAQFLIQEIDLIKTGQPLVGIETLPPDLRDFV EEDNQRFEKELEEWDAQLAQKALQEKLLAS QKLRESETSVTTAQAAGDPKYLEQPSRSDFSK HLKEETIQIITKASHEHEDKSPETVLQSAIKLE YARLVKLAQEDTPPETDYRLHHVVVYFIQNQ APKKIIEKTLLEQFGDRNLSFDERCHNIMKVA QAKLEMIKPEEVNLEEYEEWHQDYRKFRETT MYLIIGLENFQRESYIDSLFLICAYQNNKELL SKGLYRGHDEELISHYRRECLLKLNEQAAELF ESGEDREVNNGLIIMNEFIVPFLPLLLVDEMEE KDILAVEDMRNRWCSYLGQEMEPHLQEKLT DFLPKLLDCSMEIKSFHEPPKLPSYSTHELCER FARIMLSLSRTPADGR
98	1448	A	1304	118	453	SGPSSRAIYLHRKEYSONLTSEPTLLQHRVEH LMTCKQGSQRVQGPEDALQKLFEMDAHGRV WSQDLILQVRDGWLQLLDIETKEELDSYRLD SIQAMNVALNTCSYNSILS
99	1449	A	1306	3	1660	CGYFCHTTCAPQAPPCPVPPDLLRTALGVHPE TGTGTAYEGFLSVPRPSGVRRGWQRVFAALS

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  DSRLLLFDAPDLRLSPPSGALLQVLDLRDPQF SATPVLASDVIHAQSRDLPRIFRVTTSQLAVPP TTCTVLLLAESEGERERWLQVLGELQRLLLD ARPRPRPVYTLKEAYDNGLPLLPHTLCAAILD QDRLALGTEEGLFVIHLRSNDIFQVGECRRVQ QLTLSPSAGLLVVLCGRGPSVRLFALAELENI EVEVPKIPESRGCQVLAAGSILQARTPVLCVA VKRQVLCYQLGPGPGPWQRRIRELQAPATVQ SLGLLGDRLCVGAAGGFALYPLLNEAAPLAL
						GAGLVPEELPPSRGGLGEALGAVELSLSEFLL LFTTAGIYVDGAGRKSRGHELLWPAAPMGW GYAAPYLTVFSENSIDVFDVRRAEWVQTVPL KK\VRPLNPEGSLFLYGTEKVRLTYLRNQLAE KDEFDIPDLTDNSRRQLFRTKSKRRFFFRVSE EQQKQQRREMLKDPFVRSKLISPPTNFNHLV HVGPANGRPGARDKSP
100	1450	A	1318	918	190	SLCVPGPVDTGTFAVMSVMVGSVTESLAPQA LNDSMINETARDAARVQVASTLSVLVGLFQV GLGLIHFGFVVTYLSEPLVRGYTTAAAVQVF VSQLKYVFGLHLSSHSGPLSLIYTVLEVCWKL PQSKVGTVVTAAVAGVVLVVVKLLNDKLQQ QLPMPIPGELLTLIGATGISYGMGLKHRFEAG\ PPVAPNTQLFSKLVGSAFTIAVVGFAIAISLGK IFALRHGYRVDSNQVWVMRDV
101	1451	A	1353	220	445	DWPDLFTYPLIGSPKCFQSARPE\RMYRRTVR SSHGNHALQEVLPRSGHGTEFTKQKHLEAAD HGHPPARMSIFSR
102	1452	A	1363	542	2	AHLLMLNLAL\TDLL\YLTSLPFLIHYYASGEN WIFGDFMCKFIRFSFHFNLYSSILFLTCFSIFRY CVIIHPMSCFSIHKTRCAVVACAVVWIISLVA VIPMTFLITSTNRTNRSACLDLTSSDELNTIKW YNLILTA\LLCLPLVIVTLCYTTIIHTLTHGHAN \DSCLKQKARRLTILLL
103	1453	A	1371	2	410	CHSTESSSDFILPGDYLLGGLCPLHSGCLQVC SFNEHGYHLFQAMRLAVEEINNSTALLPNITL GYQLYDVCSDSANVYATLRVLSLPGQHHIEL QGDLLHYSPTVLAVIGPDSTNRAATTAALLSP FLVPMLLEQ
104	1454	A	1376	3	432	NSRVEDRS/NMSLWTQNITVCPVRNVTRDGG FGPWSPWQPCEHLDGDNSGSCLCRARSCDSP RPRCGGLDCLGPAIHIANCSRNGAWTPWSSW ALCSTSCGIGFQVRQRSCSNPAPRHGGRICVG KSREERFCNENTPCPVPIF
105	1455	A	1379	2	396	GLGLLYLIFAAVEGVMRVIGGSNHLAVVLDD IILAVIDSIFVWFIFISLAQTMKTLRLRKNTVKF SLYRHFKNTLIFAVLASIVFMGWTTKTFRIAK CQSDWMERWVDDAFWSFLF\SLILIVIMFLW RPSA
106	1456	A	1383	1	432	EDGHGGWSSRCLVDHAEEGHREPWKRLCIW QRGGHEIRFAFYFPGHPLLSPQICLAPETPPRG CPPVSSLHFISLQ/RLPRDCQELFQVGERQSGL FEIQPQGSPPFLVNCKMTSGTFWTCRTDSRVF QNANPSNAAHSEDQPTP
107	1457	A	1386	719	558	FFFVTRSHSVAQAECSGVFTAHRSLDLVGSSN YPALSLQSSWDHRHTWLIFAFL
108	1458	A	1397	631	2	RVAISLLCAAIFISFMVQSAGKRWPTGVMLM VVVLFAFLYSWPIQALLPTYLKTDLAYNPHT VANVLSFSGFGAAVGCCV/GGFLGDWLGTRK AYVCSLLASQLLIIPVFAIGGANVWVLGLLLF

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  FQQMLGQGIAGILPKLIGGYFDTDQRAAGLG
109	1459	A	1402	15	387	FTYNVGALGGALAPIIGALIAQRLDLGTALAS LSFSLTFVVILRNRRPGKSLVR VLVALPDT\VTSETVVTEVLGHRVTLPCLYSS
					;	WSHNSNSMCWGKDQCPYSGCKEALIRTDGM RVTSRKSAKYRLQGTIPRGDVSLTILNPSESDS GVYCCRIEVPGWFNDVKINVRLNLQRASTT
110	1460	A	1421	3	350	HEDLSSLLTRGSGNQERERQLKKLISLRDWM LAELAFPVGVLATCA*SLLSC*YCVILFPCSCF FFHSPDALFSLLLLSCYFPSYCFFYYLFFSSSPL CLLLASSPFPLFILLASL
111	1461	A	1426		344	FTSTMTKPFEKESEQPA*ATLAFGAQTSTTAD QCALKPDLSYLNNSSSSSSTPATSAGGGIFGSS TSSSNPPVATFVFGQSSDPVSSYGFVNTAESST SDSLLFSQDSKLATTS
112	1462	A	1434	46	372	TTSWTTSCTRSCT*SGASSGPGWTPRTTWWR SRRSSQRTCSRACSGAWSRTW*RSS*TSSSSC STSCSSSSRSCGRPGGPLGARGVHITSCLNSC MSSSTTSSTTSTF
113	1463	A	1439	3	292	HEDIMTHYDRLVDE*ALNAGKQRYEKMISG MYLGEIVRNILIDFTKKGFLLRGQISEMLKTR GIFLTFLLSNFLIVCVLLFYVSFYLFQSCINFVL
114	1464	A	1463	1	396	KQQAVPEPHSSTTTPQEQEQNWYGQDLLNLQ QRTKVHLPGHKTGPAVAKDTPEPVKKEFTVP ATSQGP*SPFSEEPPLPPSNEEVPPTLPP*EPQS EDP*KNA*LKQMHAATTHWQQHQQHQVGC QYHGIMQ
115	1465	A	1464	291	2	AGSYPSMVWSCHWGVTQKRRAL*VYSFEEG GRRKCGQYWPLEKDSRIRFGFLTVSNLGVEN MNHYKKSTLEILNPEVNPGFFFLTLWKQGEN NYCN
116	1466	A	1465	667	337	LPPQRPA*TDSYSTCNVSSGFLAGQSHNIHLQ YWTKYQVWEWLQHFLDTNQLDANCIPFQEF DINGEHLCSMSLQEFTRAAGTAGQLLYSNLQ HLKWNGDSLFLCLSLPC
117	1467	A	1479	1	381	GTSGGPKRVLVTERFPWQNPLPVNRGQAQR VLGPSNSFQRVPLQAQKLVSSHKPGQNQKHK QLQATSVPHPVCMPLNNTQKSKQPLPSAPEN NPEEELASDPNNEESL*RPWALEDFEIGRPLG KGK
118	1468	A	1485	3	385	TYLWL*GNPPFYEKNDGGLFELILRAKDEFNS PYWDDMSDSAKHFIRPLTGRDP*KPFPCDQPL QHPWIEGHTCLDNNIHQAASEPINNNFAESKR NLAFLATGVVRHMRKLFMGANLEGPGPTVS H
119	1469	A	1486	1	398	GTTSKHH*LARSLIRGPFDHDLKPNAATRDQL NIIVSYPPTKQLTYEEQDLGWKFRYYLTNQE KALTKFLKWVNWDLPQEAKQALELLGKWK PMDVKDSLELLSSHYTNPTVRRYAVARLRQA DDEDLLMYL
120	1470	A	1497	3	999	MGESPAV*GYFVLAGMNSAGLSFGGGAGKY LAEWMVHGYPSENVWELDLKRFGALQSSRT FLRHRVMEVMPLMYDLKVPHWDFQTGRQL RTSPLYDRLDAQGARWMEKHGFERPKYFVP PDKDLLALEQSKTFYKPDWFDIVESEVKCCK EAVCVIDMSSFTEFEITSTGDQALEVLQYLFS NDLDVPVGHIVHTGMLNEGGGYENDCSIARL NKRSFFMISPTDQQVHCWAWLKKHMPKDSN LLLEDVTWKYTALNLIGPRAVDVLSELSYAP

SEQ ID NO: of	SEQ ID NO: of peptide	Met hod	SEQ ID NO:	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid,
nucl- eotide	seq-	1	in USSN	nucleotide location	location corresponding	F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine.
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence			914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
				amino acid residue of	of peptide	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon,
	}	}		peptide	sequence	/=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion
						MTPDHFPSLFCKEMSVGYANGIRVMSMTHT
						GEPGFMLYIPIEYRWGFTMLSTLVSNS
121	1471	A	1498	3	306	AQFLLVGWDHIL*LIVL*TNLTELGRTTCDQN
						WPNSPDVLNHGCFYMQCLSKDCTIGYVSRE
	}		1	}		MLVAHTHTVEEHTGTHLQYVSWPDHSVPDD SSDFVEFEN
122	1472	A	1533	121	329	LGLFSFVWTEVLEEPKDFSCETEDFKTLHCT
122	1472	Α.	1555	121	329	WDPGTDTALGWSKQPSQSYTLFES*VGSGYII
						DNFFLA
123	1473	A	1547	111	408	DARTTWKPRNGSSGIWPGDGAK*PPAVEQAE
1	}	1				RGHVEMIEKLTFLNLHTSEKDKGGNTALHLA
		Į				AKHGHSPAVQVLLAQWQDINEMNEKQQTPL
101	1454		1		745	HVAADRG
124	1474	Α	1555	1	745	MTFDDDDKNTYGVALVWKKFQTQSLRLSDL HRKSHLWRGIVSITLIEGRDLKAMDSNGLSDP
						YVKFRLGHQKYKSKIMPKTLNPQWREQFDF
						HLYEERGGVIDITAWDKDAGKRDDFIGRCQV
		ļ		ļ		DLSALSREQTHKLELQLEEGEGHLVLLVTLT
		ļ				ASATVSISDLSVNSLEDQKEREEILKRYSPLRI
		}		ļ		FHNLKDVGFLQVKVIRAEGLMAADVTGKSD
						PFCVVELNNDRLLTHTVYKNLNPEWNKVFTL
Į						*VALVWKKFQTQSLRLSDLHRKSHLWRGIVS
						ITLIEGRDLKAMDSNGLSDPYVKFRLGHQKY   KSKIMPKTLNPOWREOFDFHLYEERGGVIDIT
		}	ļ			AWDKDAGKRDDFIGRCQVDLSALSREQTHK
			}			LELQLEEGEGHLVLLVTLTASATVSISDLSVN
	·					SLEDQKEREEILKRYSPLRIFHNLKDVGFLQV
		}				KVIRAEGLMAADVTGKSDPFCVVELNNDRLL
						THTVYKNLNPEWNKVFTL
125	1475	Α	1556	57	509	GGPAPNSRYAEP*KNSLAMT*AHADCENYVA
					1	CGGLDNICSIYNLKTREGNVRVSRELPGHTGY
		}				LSCCRFLDDSQIVTSSGDTTCALWDIETAQQT TTFTGHSGDVMSLSLSPDMRTFVSGACDASS
						KLWDIRDGMCRQSFTGHVSDINAVS
126	1476	A	1592	3	178	KSEKSCVSSLAHFGTSCQRDYDAMVKLVETL
						EMLPTCDLADQHNIKFHYAFALNR*ER
127	1477	A	1612	1	497	TESPLLVRPYLPYITKSELHAIMTAGFSTIAGS
						VLGAYISFGVPSSHLLTASVMSAPASLAAAKL
						FWPETEKPKITLKNAMKMESGDSGNLL*AAT
						QGASSSISLVANIAVNLIAFLALLSFMNSALA WVGNMFDYPQLSFELICSYIFMPFSFMMGVE
						WPDSFM
128	1478	A	1619	286	486	CCMNSKAQESVFKNVLCNPPALSEMPDVKA
						EDEVDFRASSISEEVAVGSIAATLKMKQGPM
						TQAINR
129	1479	A	1627	1	395	PTRGALRYWIFGRFLCNIWAAVDVRCCTATI
			}	}		MGLCIISIDRYVGVSYPLRYPTIVTQRRGLMA
						LLCVWALSLVIYIGPLLGWRHPAPEDETICQI
{			}			NEEPGYVLFSTPGSFYLPLAIMLVMN*RVYRV AKTE
130	1480	Ā	1638	2	466	DPRVRTKIVNRKTTIYEIQDKTGSMAVVGKG
130	1400	<b>'`</b>	1050	-	-100	ECHNIPCEKGDKLRLFCFRLRKRENMSKLMS
						EMHSFIQIQKNTNQRSHDSRSMALPQEQSQHP
						KPSEASTTLPESHLKTPQMPPTTPSSSSFTKVT
						KDKDIK*LLFNLYSSVEILPEVLHLKT
131	1481	A	1651	607	3	LAEGGDVFDCVLNGGPLPESRAKALFRQMVE
						AIRYCHGCGVAHRDLKCENALLQGFNLKLTD
						FGFAKVLPKSHRELSQTFCGSTAYAAPEVLQ GIPHDSKKGDVWSMGVVLYVMLCASLPFDD
L		<u> </u>				CIT I DOKTOD A MOMO A APT AMERICAPELEDD

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  TDIPKMLWQQQKGVSFPTHLSISADCQDLLK
132	1482	A	1656	150	48	RLLEPDMILRPSIEEVSWHPWLAST**KQWQV LSNKVGGESKPKKKK LVAKSLLYCGCLFFLLQLAKNVGNNSFNDIM EANLTSPSPKPTPSSDM*VFLIY*TYFGAWHV
133	1483	A	1660	3	406	VDAQ RKHIKLLIQKLSDVP*ECQNNQL*KLTEICEKE KKEFKKKMDDQRPEKITEA*SKDKSPMEEEK TEMIRSYIQEVGRYIKRLEEAQSKRLEKLREK HKEIRQPILDEKPKGEGSSSFLSETCHEDTSWF PNFTP
134	1484	A	1666	1276	466	PGSTHASARITIY*L*IILSNATEVDNNFSKPPP FFPAGAPPASSSSSSSSSSSPTVSTAPPLIPPPGF PPPPGAPPPSLIPTIESGHSSGYDSRSARAFPYG NVAFPHLPGSAPSWPSLVDTSKQWDYYARSS SSSSSSSSSSSPRDRDRER*RTRERERERDHS PTPSVFNSDEERYRYREYAERGYERHRASRE KEERHRERRHREKEETRHKSSRSNSRRRHESE EGDSHRRHKHKKSKRSKEGKEAGSEPAPEQE STEATPAE
135	1485	A	1673	1	417	PTRPVNSSQAFALVYYTLGALGGNLIAHMGL GYRYWAGIGVLQSCESALTHYRLVANHVAS DISLTGGSVVQRIRLPDEVENPGMNSGMLQE DLIQYYQFLAEKGDVQAQVGLGQLHLHGGR GV*QNHQRAFDYFNLAA
136	1486	A	1678	525	9	ANTSLSSAAVSAVSPPPCRTSTATTLPPPMPSF FCVFPSPSMSPSPSEFLSCIASVSRVHSLSSSSS GSSSTASSLNFSAIMGSSSATASWVLSTASTPP CPSALPSSPAQES*SLAASSSAWPVAGISPSGA CTFPAGSASGAAKAPSPSWRCPSFRALFSLLD SSSLSL
137	1487	A	1680			AHRDEIQRKFDALRNSCTVITDLEEQLNQLTE DNAELNNQNFYLSKQLDEASGANDEIVQLRS EVDHLRREITEREMQLTSQKQTMEALKTTCT MLEEQVMDLEALNDELLEKERQWEAWRSVL GDEKSQFECRVRELQRMLDTEKQSRARADQ RITESRQVVELAVKEHKAEILALQQALKEQK LKAESLSDKLNDLEKKHAMLEMNARSLQQK LETERELKQRLLEEQAKLQQMDLQKNHIFR LTQGLQEALDRADLLKTERSDLEYQLENIQV LYSHEKVKMEGTISQTKLIDFLQAKMDQPA KKKKVPLQYNELKLALEKEKARCAELEEALQ KTRIELRSAREEAAHRKATDHPHPSTPATARQ QIAMSAIVRSPEHQPSAMSLLAPPSSRRKESST PEEFSRRLKERMHHNIPHRFNVGLNMRATKC AVCLDTVHFGRQASKCLECQVMCHPKCSTC LPATCGLPAEYVTHFTEAFCRDKMNSPGLQT KEPSSSLHLEGWMKVPRNNKRGQQGWDRK YTVLEGSKVLIYDNEAREAGQRPVEEFELCLP DGDVSIHGAVGASELANTAKADVPYILKMES HPHTTCWPGRTLYLLAPSFPDKQRWVTALES VVAGGRVSREKAEADAKLLGNSLLKLEGDD RLDMNCTLPFSDQVVLVGTEEGLYALNVLK NSLTHVPGIGAVFQIYIIKDLEKLLMIAGEERA LCLVDVKKVKQSLAQSHLPAQPDISPNIFEAV KGCHLFGAGKIENGLCICAAMPSKVVILRYN ENLSKYCIRKEIETSEPCSCIHFTNYSILIGTNK FYEIDMKQYTLEEFLDKNDHSLAPAVFAASS NSFPVSIVQVNSAGQREEYLLCFHEFGVFVDS

SEQ ID NO: of	SEQ ID NO: of	Met hod	SEQ ID NO:	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid,
nucl- eotide	peptide seq-		in USSN	nucleotide location	location corresponding	F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence		1	914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
				amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
ĺ				residue of peptide	sequence	Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible
		}		sequence		nucleotide insertion
						YGRRSRTDDLKWSRLPLAFAYREPYLFVTHF
						NSLEVIEIQARSSAGTPARAYLDIPNPRYLGPA
1						ISSGAIYLASSYQDKLRVICCKGNLVKESGTE HHRGPSTSRR*PASPLPQYQGQRAFLQGRRK
138	1488	A	1686	2	526	GRPQGPAPGAGSPPESGPGLWAALGCSLVWV
						PLCCLGGAAGRL*ARSGKSGLRRRRAHAGPP
	]	}		ļ	j	PGGPCNSCP*CSAPESGGRGPLPGPGTGGVCS
						CWTRGCQTTARTAAAAAAPGPAGRRPPGGA PQNGSCAASASQEAAAPPPMCPPGRRWAVAS
	ł					PPETRCPAAPGTRCRRLEAA
139	1489	A	1693	3	376	LPSMSNCTSCFRLQSRTES*IRQAGHLLGRNE
	Ì					FIETKALGCAWFSLCYYLVLYFESSHKVDFVF
						IV*CFSTPPGAQMTIMSQACAERCNIMRLVDR
140	1490	A	1704	3	376	RWAGIAKGVGTQKIIGRVHLGEQKALGL ERTNKFIKELIMDGKNLIAATKSLSVAQRKFA
140	1430	A	1704	, ,	370	HSLRDFKFEFIGDAVTDDERCIDASLREFSNFL
						KNLEEQREIMVS*EGCKLISQLSRGKKIWIWK
						LVLVEVVKHLSLGTVVHCNGKMRFPEP
141	1491	A	1743	1	362	LITNKVFVARELSCLDVHLDSTGSTAVVADQ
					1	DKLELELVLKGSYEDTQTSFLGTASAFRFHY MAAL*TELSGRLRSSKSNGWNGDNSTGYLTV
	İ				1	PLRPLTIVKEVTMDVPAPNVRGLNWMG
142	1492	A	1769	1	406	NNPSTLPRGS*PMSPRTTMGRRRQRRREHKSS
						LSLASSTVGPGGQIVHTETTEVVLCGDPLSGF
						GLQLQGGIFATETLSSPPLVCFIEPDSPAERCG LLQVGDRVLSINGIATEDGTMEEANQLLRDA
					,	ALAHKVV
143	1493	A	1789	1	447	QMLRNGGDQNTVPDYHFADRIRELL*PTEDQ
		1				KNCIP*DTYLRPSALGNIVEEVTHPCSPGPCPA
						NELCEVNRKGCTSGDPCLPYFCVQGCKLGQA
	ĺ	ļ				SDFIARQGTLIQVPSSAGEVECYKICSCGQSGL LENCMEMHCMDLPTDTSALVR
144	1494	A	1814	1	404	PGRRFRPRLSQAGTDSGS*VFPDSFPSAPAEPL
						PYFLQEPQDAYIVKNKPVELRCRAFPATQIYF
	1					KCNGEWVSQNDHVTQEGLDEATGLRVREVH
						IEVSRQQVEELFGLEDYWCQCVAWSSAGTTK   SRRAYVRI
145	1495	A	1827	26	448	XVEEKHADTWRSXCLSDFFFHAAKXLCXE*N
		1				CGDAISLSVGDHFGKGNGLTWAEKFQCEGSE
	]	1				THLALCPIVQHPEDTCIHSREVGVVCSRYTDV
	ļ					RLVNGKSQCDGQVEINVLGHWGSLCDTHWD PEDARVLCRQLNCGTAL
146	1496	A	1828	574	333	QHEGGDLRRRQLGEIQLTVRYVCLRAASAC*
			-3-5			SMAAET*HHVPASGADPYVRVYLLPERKWA
						CRKKTSVKRKTLEPLFDET
147	1497	A	1855	1	372	ERLVLTSEHCLVLTLFWPSWTYHTLLLSRQH
		Į				VRRLPKLTHAEHDHLASIMNKLLTNYDNLFE TSVTYSMG*HGAPTGSEAGANWNH**LHAH
						YYPPLLRSDTVRKFMVGSQMLAQAQRDLTPE
		ļ				Q
148	1498	A	1879	568	7	LLSALDDKGGTQPSASFSNAPTIVCVTACPAG
}	<b>!</b>	l				IAHTYMAAEYLEKAGRKLGVNVYVEKQGAN
[		1				GIEGRLTADQLNSATACIFAAEVAIKESERFN   GIPALSVPVAEPIRHAEALMQQALTLKRSDET
	[	ĺ	[			RTVQQDTQPVKSVKTELKQALLSGISFAVPLI
						VAGGTQVA*AV*RQGISSLHDVQVRTWNS
149	1499	A	1880	611	24	GLNSENALSNEAMERGWQCLRLFAERLQDIP
		1				PSQIRVVATATLRLAVNAGDFIAKAQEILGCP VQVISGEEEARLIYQGVAHTTGGADQRLVVD
L	L			L	L	- 4 - 12 ODDE HOLL GO - ALLI TOURD QUE VVD

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  IGGASTELVTGTGAQTT*LFSLSMGCVTWLER YFADRNLGQENFDAAQKAAREVLRPVADEL RYHSWKEVRGASVTVQALQEIMMAQGMDE RITMEIWPVD  GRVDFFHTDYRPLIRDSNNYVLDEQTQQAPH
						LMPPPFLVDVDGNPHPTKYQRLVPGRENSAD EHLIPQLGYVATSDGEVIEQIISLQTNDNDERS PESSILDGMIRQLQQQQDQRMGADQDTIPRG LSNGEETPRRGFRRLSLDIQSPPNIGLRRSGQV EGVRQMHQNAPRSQIATERDLQAWKRRVVV PEVPLGIFRKLEDFRLEKGEEERNLYIIGRKRK TLQLSHKSDSVGLVSQSRPRTCRRKYP
151	1501	A	1900	141	785	GKTIQIQTTMQNKYKTVQKQYKTIPKNKKA MEMQIKKQFQDTCKVQTKQYKALKNHQLEV TPKNEHKTILKTLKDEQTRKLAILAEQYEQSI NEMMASQALRLDEAQEAECQALRLQLQQEM ELLNAYQSKIKMQTEAQHERELQKLEQRVSL RRAHLEQKIEEELAALQKERSERIKNLLERQE REIETFDMESLRMGFGNLVTLDFPKEDYR
152	1502	A	1915	2	377	LVRLLDTQRDGLQNYEALLGLTNLSGRSDKL RQKIFKERALPDIENYMFENHDQLRQAATEC MCNMVLHKEVQERFLADGNDRLKLVVLLCG EDDDKVQNAAAGALAMLTAAHKKLCLKMT QVTT
153	1503	A	1921	1	237	AYQSLRLEYLQIPPVSRAYTTACVLTSAAVQL ELITPFQLYFIPELIFKHFQIWRLITNFLFFVPFG FNFLLYMIFLYT
154	1504	A	1928	2	354	EMVEGGEGKMCINTEWGGFGDNGCIDDIRTR YDTEVDEGSLNPGKQRYEKMTSGMYLGEIV RQILIDLTKQGLLFRGQISERLRTRGIFETKFLS QIESDRLALLQVRRILQQLGLD
155	1505	A	1929	2	369	TEIAKIKMEAKKKYEKELTMFQNDFEKACQA KSEALVLREKSTLERIHKHQEIETKEIYAQRQ LLLKDMDLLRGREAELKQRVEAFESYQLELK DDYIIRTYRLIEDDRINIQISGHWQESP
156	1506	A	1935	1	270	VTRKLPIFIVDAFTARAFRGSPAADCLLENEL DEDMHQKIAREMNLSETAFIRKLHPTDNFAQ RSCFGLIWFTPTTDLQILTSSILPSIL
157	1507	A	1936	584	305	ESKVNNEKFRTKSPKPAESPQSATKQLDQPTA AYEYYDAGNHWCKDCNTICGTMFDFFTHMH NKKHTQGQFQKSSDFQKEELQQTFLPPERQG
158	1508	A	1939	1	423	TTHRLNVTAEPPCTSMPIYWMPDVPHRCTTA NTCPVDLTDYCAQNGFYCLVYGFLPYGSLED RLHCQTQACPPLSWPQRLDILLGTARAIQFLH QDSPSLIHGDIKSSNVLLDERLTPKLGDFGLA RFSRFAGSSPIQSSM
159	1509	A	1974	3	401	HTSTARLLLHRGAGKEAVTSDGYTALHLAAR NGHLATVKLLVEEKADVLARGPLNQTALHL AAAHGHSEVVEELVSADVIDLFDEQGLSALH LAAQGRHAQTVETLLRHGAHINLQSLKFQGG HGPAATLLR
160	1510	A	1982	2	417	KFLKDLEKQYNKEEPHLSEIGSCFLQNQEGFA IYSEYCNNHPGACLELANLMKQGKYRHFFEA CRLLQQMIDIAIDGFLLTPVQKICKYPLQLAEL LKYTTQEHGDYSNIKAAYEAMKNVACLINER KRKLESIDKIA
161	1511	A	1984	4	770	RETGSVSLSPSGLEGAESYAVSPILYSSPDVKE LWLETLQGQRHSHTGVKSTPGQSAAILMKLR SSHNASKTLNANNMETLIECQSEGDIKEHPLL

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162	1512	Ā	1986	864	501	RKLLSSDLFEEWMGALEMQDEEDRIEALK LLNSGLFSAPDGSNLEMRLTRGGNMCSGRIEI KFQGRWGTVCDDNFNIDHASVICRQLECGSA VSFSGSSNFGEGSGPIWFDDLICNGNESALWN CKHQGWGKHNCDHAEDAGVICSSKD
163	1513	A	2001	419	187	AVDLSIDESSLTGETTPCSKVTAPQPAATNGD LASRSNIAFMGTLVRCGKAKGVVIGTGENSE FGDIINLSTFVVHS
164	1514	A	2012	284	597	SLLCLFPGTSTVVCKPIVIETQLYVIVAQLFGG SHIYKRDSFANKFIKIQAIEILKIRKPNDIETFKI ENNWYFVVADSSKAGFTTIYKWERETGFYSH QSFTR
165	1515	A	2013	2	403	EDPEELGHFYDYPMALFSTFELFLTIIDGPANY NVDLPFMYSITYAAFAIIATLLMLNLLIAMMG DTHWRVAHERDELWRAQIVATTVMLERKLP RCLWPRSGICGREYGLGDRWILRVEDRQDLN RQRIQRYA
166	1516	A	2019	2	927	CCQREGLGLKAVVQILLSHGRNGLPGEPASS QGLSAASSTPVFHLALQIDSAPDNIDWVEMLF NKNMVTERLQNVMVLEQCFSDSSSLYRFLTY SYLLAFNVWLLLAPVTLCYDWQVGSIPLVETI WDMRNLATIFLAVVMALLSLHCLAAFKRLE HKEVLVGLLFLVFPFIPASNLFFRVGFVVAER VLYMPSMGYCILFVHGLSKLCTWLNRCGATT LIVSTVLLLLLFSWKTVKQNEIWLSRESLFRS GVQTLPHNAKVHYNYANFLKDQGRNKEAIY HYRTALNNNKAWDYLCWRFRKTLTDLP
167	1517	A	2025	696	71	AAASAASSLTVTLGRLASACSHSILRPSGPGA ASLWSASRRFNSQSTSYLPGYVPKTSLSSPPW PEVVLPDPVEETRHHAEVVKKVNEMIVTGQY GRLFAVVHFASRQWKVTSEDLILIGNELDLA CGERIRLEKVLLVGADNFTLLGKPLLGKDLV RVEATVIEKTESWPRIIMRFRKRKNFKKKRIV TTPOTVLRINSIEIAPCLL
168	1518	A	2046	2	366	HLQVAARVFMPLQAVDSAPKPLKGQAQAPQ RLQGAARVFMPLQAQVKAKASKPLQMQIKA PPRLRRAARVLMPLQAQVRAPRLLQVQSQVS KKQQAQTQTSEPQDLDQVPEEFQGQDQVLR
169	1519	A	2049	1	945	QNLEDREVLNGVQTELLTSPRTKDTLSDMTR TVEISGEGGPLGIHVVPFFSSLSGRILGLFIRGI EDNSRSKREGLFHENECIVKINNVDLVDKTFA QAQDVFRQAMKSPSVLLHVLPPQNREQYEKS VIGSLNIFGNNDGVLKTKVPPPVHGKSGLKTA NLTGTDSPETDASASLQQNKSPRVPRLGGKPS SPSLSPLMGFGSNKNAKKIKIDLKKGPEGLGF TVVTRDSSIHGPGPIFVKNILPKGAAIKDGRLQ SGDRILEVNGRDVTGRTQEELVAMLRSTKQG ETASLVIARQEGHFLPRELVMFRSQSH
170	1520	A	2050	363	1	PVATHLTKILNSDEHAVVISSAKTLCETVKDF VAKVEKTYDKTLENAVVADAVASKCSVLNE KLEQLLQALHTDSQAAPVLPGLSPLIVEEDAV ESSSEESLGESKEQLGDDVTKPSSQKA
171	1521	A	2055	139	675	IPSRPWLGRITGLDPAGPLFNGKPHQDRLDPS DAQFVDVIHSDTDALGYKEPLGNIDFYPNGG LDQPGCPKTILGGFQYFKCDHQRSVYLYLSSL

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172	1522	A	2056	3	361	SCPLLGYYADNWKDHLRGKDPPMTKAFFDT AEESPFCMYHYFVDIITWNKNVR LIQHKSAVEYAQSHLSLVSMCKESHKCSEPK
172	1322	A	2030	3	301	MEWKVKIRSDGTRYITKRPVRDRILKERALKI KEERSGLTTDDDTMSEMKMGRYWSKEERKQ HLVRGKEQRRRREFMMRIRLKCLKES
173	1523	A	2060	1	387	GTRILSMQIPFVGFQPIRTSEHMAAAGVFALL QAYAFLQYLRDRLTKQEFQTLFFLGVSLAAG AVFLSVIYLTYTGYIAPWSGRFYSLWDTGYA KIHIPIIASVSEHQPTTWVSFFFDLHILGCTFPA G
174	1524	A	2071	74	443	LLMGPKAKKSGSKKKKVTKAERLKLLQEEEE RRLKEEEEARLKYEKEEMERLEIQRIEKEKW HRLEAKDLERRNEELEELYLLERCFPEAEKLK QETKLLSQWKHYIQCDGSPDPSVAQEMNT
175	1525	A	2083	139	486	AALTWSQPQEFWPMEMQPIVTDMVTVHWV AESSTVGWLCALFRVTHVGVGATGHGVVCG RRVLCGLPLPSPAPMPIMSLPEGESRKEREVQ RLQFPYLEPGHELPATTLLAFLAAV
176	1526	A	2092	3	587	EGSVNFKFGVLFAKDGQLTDDEMFSNEIGSEP FQKFLNLLGDTITLKGWTGYRGGLDTKNDTT GIHSVYTVYQGHEIMFHVSTMLPYSKENKQQ VERKRHIGNDIVTIVFQEGEESSPAFKPSMIRS HFTHIFALVRYNQQNDNYRLKIFSEESVPLFG PPLPTPPVFTDHQEFRDFLLVKLINGEKATLET PCI
177	1527	A	2103	44	427	GKGQVSLEGRPHRGPLCLGSWWPGSRVPGC CDGAWLAWACWVFGNDFPSPASAACSALLG CSVSTACLCVPLCSGSPLAPFRRTAALQEGLR RAVSVPLTLAETVASLWPALQELARCGNLAC RSDLO
178	1528	A	2104	2	409	ALQSTLGAVWLGLLLNSLWKVAESKDQVFQ PSTAASSEGAVVEIFCNHSVSNAYNFFWYLHF PGCAPRLLVKGSKPSQQGRYNMTYERFSSSL LILQVREADAAVYYCAVEVPNTDKLIFGTGT RLQVFPNIQNPD
179	1529	A	2111	1	312	PTRSSTRPPSLFVHASAKGGEKEEGDDGHYL MRTESHTGLKKGGNANLVFMLKRNTEPKKG SYHFDLERLRAAHILFEREQEHLAPGGISMPL PPPLPLACLG
180	1530	A	2116	3	366	TSIKRAIETTDVTRSFGWDSSEAWQQHDVQE LCRVMFDALEQKWKQTEQADLINELYQGKL KDYVRSLECGYEGWRIDTYLDIPLVIRPYGSS QAFASVVCTFHLTACVSLHRIHNSTVV
181	1531	A	2117	2	386	YGLGAHFGRLFIQAGINENDFYDGAWCAGR NDLQQWIEVDARRLTRFTGVITQGRNSLWLS DWVTSYKVMVSNDSHTWVTGKNGSGDMIFE GNSEKEIPVLNELPVPMVARYIRINPQSWFDN GSICI
182	1532	A	2123	1	493	RTKTDVYILNLAVADLLLLFTLPFWAVNAVH GWVLGKIMCKITSALYTLNFVSGMQFLACISI DRYVAVTKVPSQSGVGKPCWIICFCVWMAAI LLSIPQLVFYTVNDNARCIPIFPRYLGTSMKAL IQMLEICIGFVVPFLIMGVCYFITARTLMKMP NIKIS
183	1533	A	2140	3	561	RQAWHEAFKVRKEILTVICCLLAFCIGLIFVQ RSGNYFVTMFDDYSATLPLLIVVILENIAVCF VYGIDKFMEDLKDMLGFAPSRYYYYMWKYI

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						EEFLSYPTWGLAVCASLDVFAILPVPVAFIGR RFSLIDDGAGPFCSAAYTTTGCRTPYL
184	1534	A	2145	3	538	HELTVAAADRGQPPQSSVVPVTVTVLDVND NPPVFTRASYRVTVPEDTPVGAELLHVEASD ADPGPHGLVRFTVSSGDPSGLFELDESSGTLR LAHALDCETQARHQLVVQAADPAGAHFALA PVTIEVQDVNDHGPAFPLNLLSTSVAENQPPG TLVTTLHAIDGDAGAFGRLRYHL
185	1535	A	2151	2	671	LDKLLDRMENYNIFNEYILKQVAATYIKLGW PKNNFNGSLVQASYQHEELRREVIMLACSFG NKHCHQQASTLISDWISSNRNRIPLNVRDIVY CTGVSLLDEDVWEFIWMKFHSTTAVSEKKIL LEALTCSDDRNLLNRLLNLSLNSEVVLDQDAI DVIIHVARNPHGRDLAWKFFRDKWKILNTRI RQKTLEFDFAEPLILAFPIILYTAIDNPPLVREH E
186	1536	A	2153	2	400	GPMCDKHSAFAEKFHAGFIDYIVHPLWETWA HLALPDAQDILYTLEDNRNWVDSMIPQSPSPP LDEQNRDWQGLLENLHVELTLDEEDSEGPEK EGEGQTYFTSSKTLCGIVPQNTDSLGETGIHIC AHDKSP
187	1537	A	2158	227	442	FNCFRVASDSFLENSSLLIMILPLRNATQEFIIR PGAVAYTCNPSTLGGWGGWITRSGVRDQPG OHGGTPS
188	1538	A	2167	3	486	AHLGGAWLTORSLGSWAAPGPARAAKEVVA CIPQNQKMNIWRMKTSKHLQLLSFVLGAVSP AVVVPYMMVLQENGYGVEEGIPTLLMAASS MDDILAITGFNTCLSIVFSSGCARSSGSRNSKS LRTPLGTICEGCDDSSIFSHLDHSSKWSSTYG HSGA
189	1539	A	2168	2	412	EFLSSNQITQLPNTTFRPMPNLRSVDLSYNKL QALAPDLFHGLRKLTTLHMRANAIQFVPVRIF QDCRSLKFLDIGYNQLKSLARNSFAGLFKLTE LHLEHNDLVKVNFAHFPRLISLHSLCLRRNKV AIVVSSLDW
190	1540	Α	2179	64	399	MRLNQNTLLLESFGXXRPYTSEHAPTYHQW MKADELLRWTTSEPLTLEHEYAMQRTWLED AYECTFIVLDAEKRHAQPGATEESCMVGDVN LFLTDLEDLTLGEIEVLIAEP
191	1541	A	2190	1	469	CLDRAAGIRHERNVIYINETHTRHRGWLARR LSYVLFIQERDVHKGMFATNVTENVLNSSRV QEAIAEVAAELNPDGSAQQQSKAVNKVKKK AKRILQEMVATVSPAMIRLTGWVLLKLFNSF FWNIQIHKGQLEMVKAATETNLPLLFLPVHR SH
192	1542	A	2197	26	157	PSKXGGIRLLLTGTQLYGRFGSAIAPLGDLDR DGYNGEGREEPY
193	1543	A	2236	2	383	EYFPNSIWRSLFSTMDLGDIGFYTYRILQALS YTHSKGIMHRDVKPLNILCNSPRNKVILADW GLAEFYHPMRKYSVHVATRYYKSPEILLDYE YYDYSLDIWAVGVILLELLTLKLHVFEGGDN EQ
194	1544	A	2241	105	409	RKGVGKMPTSEGRPGQERSDWVTSYKVMGS NDSHTWVTVKNGSGDMIFEGNSEKEIPVLNE LPVPMGARYIRINPQSWFDNGSICMRMEILGC PLPDPNNY
195	1545	A	2245	1	672	MGVASDWTKRIEYQPGSGSMPLFPSIHLETCD GAVSSLQIVTELQTNYIGKGCDRETYSEKSLQ

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196	1546	A	2256	1	396	ISLKPS FRGTPVSGLTNRDTLAVIRHFREPIRLKTVKP GKVINKDLRHYLSLQFQKGSIDHKLQQVIRD NLYLRTIPCTTRAPRDGEVPGVDYNFISVEQF KALEESGALLESGTYDGNFYGTPKPPAEPSPF OPDPV
197	1547	A	2259	43	594	QLAIEIGVRALLFGVFVFTEFLDPFQRVIQPEEI WLYKNPLGQSDNIPTRLMFAISFLTPLAVICV VKIIRRTDKTEIKEAFLAVSLALALNGVCTNTI KLIVGRPRPDFFYRCFPDGVMNSEMHCTGDP DLVSEGRKSFPSIHSSFAFSGLGFTTFYLAGKL HCFTESGRGKSWRLCAAILPL
198	1548	A	2275	3	404	TCTTVVVIPRMLVDFLSESKTISLPECATQMFF FLGFASNNCFIMAAMSYDRYTAIHNPLQYHT LMTRKICLQMMMASWMVGFLFSLCIIVTVFN LSLCDLNTIQHYFCDISPVVSLACNYTFYHEM AIFVLSA
199	1549	A	2315	1	375	LTQMFFIHALSAIESTILLAMAFDRYVAICHPL RHAAVLNNTVTAQIGIVAVVRGSLFFFPLPLLI KRLAFCHSNVLSHSYCVHQDVMKLAYADTL PNVVYGLTAILLVMGXDRMFISLSYFLII
200	1550	A	2334	2	409	PRVRPQQRKMSFFFKTELGEKLVTKFLFETDF SDDPMLPSPDQLKKKAPFTNKKLKAHQTPVD ILKQKAHQLASMQVQAYNGGNANPRPANNE EEEDEEDEYDYDYESLSDDNILEDRPENKSCH DQLQFEYKEEM
201	1551	A	2350	3	512	ISWEAQIAEIIQWVSDEKDARGYLQALASKM TEELEALRSSSLGSRTLDPLWKVRRSQKLDM SARLELQSALEAEIRAKQLVQEELRKVKDAN LTLESKLKDSEAKNRELLEEMEILKKKMEEK FRADTGKLMLCDSALFEYKYFSNECFYFLFD LIVTLEAPTEFQIQY
202	1552	A	2351	1	1003	PSSYSSDELSPGEPLTSPPWAPLGAPERPEHLL NRVLERLAGGATRDSAASDILLDDIVLTHSLF LPTEKFLQELHQYFVRAGGMEGPEGLGRKQA CLAMLLHFLDTYQGLLQEEEGAGHIIKDLYL LIMKDESLYQGLREDTLRLHQLVETVELKIPE ENQPPSKQVKPLFRHFRRIDSCLQTRVAFRGS DEIFCRVYMPDHSYVTIRSRLSASVQDILGSV TEKLQYSEEPAGREDSLILVAVSSSGEKVLLQ PTEDCVFTALGINSHLFACTRDSYEALVPLPE EIQVSPGDTEIHRVEPEDVANHLTAFHWELFR CVHELEFVDYVFHGE
203	1553	A	2361	2	403	NNLNCAEPLFEQNNSLNVNFNTQKKTVWLIH GYRPVGSIPLWLQNFVRILLNEEDMNVIVVD WSRGATTFIYNRAVKNTRKVAVSLSVHIKNL LKHGASLDNFHFIGGSLGAHISGFVGKIFHGQ LGRITGLDP
204	1554	A	2390	280	476	SPSLLPQCLMSLSDLSLSPAPPSHLSPRCPSPQ AGSRLGAMRRCAREMDATPMPPAPSCPSERV T
205	1555	A	2400	543	745	AAVALRDISWQQPYPMDFYAGSSLGPWTVN HGQDRRPHAPGRPARGKVQEGSARPPSAVAC EDCSCR

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206	1556	A	2406	122	485	DLSPDSREDHPQGHRRLLPKRPVRGSLMPGH THHPCPVSSTTNDTPDQIWVSVGSLRMGTGG MGANASTSPRCWDLSSGNKKWIIQVPILASIV ESRGGLLATGVGGMCACVPRNQPLTGT
207	1557	A	2409	289	418	LWTLYRHKQQVQHNHSNRLSCRPSQEDRAT HTIMVLDKENTLS
208	1558	A	2413	64	492	VQGTGXXFIAFTEAMTHFPASPVWAGMFFL MLINLGLGSMIGTMAGITTPIIDTFKVPKEMFT GGCCVFAFLVGLLFVQRSGNYFVTMFDDYSA TLPLTLIVILENIAVAWIYGTKKFMQELTEML GFRPYRFYFYMWKFVSP
209	1559	A	2417	3	877	EKERLLDEWFTLDEVPKGKLHLRLEWLTLMP NASNLDKVLTDIKADKDQANDGLSSALLILY LDSARNLPIRYKTNEPVWEENFTFFIHNPKRQ DLEVEVRDEQHQCPLGNLKVPLSQLLTSEDM TVSQRFQLGNSGPNSTIKMKIALRVLHLEKRE RPPDHQHSAQVKRPSVSKEGRKTSIKSHMSG SPGPGGSNTAPSTPVIGGSDKPGMEEKAQPPE AGPQGLHDLGRSSSSLLASPGHISVKEPTPSIA SDISLPIATQELRQRLRQLENGTTLGQSPLGQI QLTIP
210	1560	A	2422	35	456	REFAASDLEPFTPTDQPISPEAITQPSCIKRQRA AGNPGSLAATIDHKPCSAPLEPKIQASRNQRW GAVRAAESLTDIAEPASPQVHETPIDASQTQK VEPASKSRFTPELQAKVSHSRERALSTMDATP HHAQPQRGEG
211	1561	A	2431	1	764	RRYSQKLIQHTACQLLRTYPAATRIDSSNPNP LMFWLHGIQLVALNYQTDDLPLHLNAAMFE ANGGCGYVLKPPVLWDKNCPMYQKFSPLER DLDSMDPAVYSLTIVSGQNVCPSNSMGSPCIE VDVLGMPLDSCHFRTKPIHRNTLNPMWNEQF LFHVHFEDLVFLRFAVVENNSSAVTAQRIIPL KALKRGYRHLQLRNLHNEVLEISSLFINSRRM EENSSGNTMSASSMFNTEERKCLQTHRVTVH GVPG
212	1562	A	2436	1	411	GIRGTTGHLGCPINDDPSLTLTVSWVMEDKPI YIGNGTKKEDDSLTIFAVAKRDHVSDTCGAC TDLDHNLDKGYLTVLGEQATPTNRLGALPKG RANRTRDLELTYLAERIVRLTWIPGDANNRPI TDYDCQIEEHQ
213	1563	A	2445	1	1294	MSSIGCLWVSRSSQIDGLTAEKSGPEKPHGT WLMPELHPKEQILELLVLEQFLSILPEELQIWV QQHNPESGEESVTLLEDLEREFDDPGQQVPAS PQGPAVPWKDLTCLRASQESTDIHLQPLKTQ LKSWKPCLSPKSDCENSETATKEGISEEKSQG LPQEPSFRGISEHESNLVWKQGSATGEKLRSP SQGGSFSQVIFTNKSLGKRDLYDEAERCLILT TDSIMCQKVPPEERPYRCDVCGHSFKQHSSLT QHQRIHTGEKPYKCNQCGKAFSLRSYLIIHQR IHSGEKAYECSECGKAFNQSSALIRHRKIHTG EKACKCNECGKAFSQSSYLIIHQRIHTGEKPY ECNECGKTFSQSSKLIRHQRIHTGERPYECNE CGKAFRQSSELITHQRIHSGEKPYECSECGKA FSLSSNLIRHQRIHSG
214	1564	A	2461	1	615	GIPGSTISSSRNIFLEDDLAWQSLIHPDSSNTPL STRLVSVQEDAGKSPARNRSASITNLSLDRSG SPMVPSYETSVSPQANRTYVRTETTEDERKIL LDSVQLKDLWKKICHHSSGMEFQDHRYWLR THPNCIVGKELVNWLIRNGHIATRAQAIAIGQ

SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide	}	in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence	ł		914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
1			ŀ	amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
	1	1	ľ	peptide		/=possible nucleotide deletion, \=possible
	ļ		ļ	sequence		nucleotide insertion
						AMVDGRWLDCVSHHDQLFRDEYALYRPLQV
215	1565		2464	l	2022	LFSVYCQLECSKLIL
215	1565	A	2464	3	2932	GPGVRSSQDGMADVFVHLRTAWPRCSFISGQ
ŀ	1	ļ	ļ			HGPGRHGRRVCSSQDSMADVFVHLRTAWPT CSLISGOHGPGESVSYEDDDIPAPASLLHVNA
						AAPALTNPTAPVLCTAPNNTAQKEKVPSGMR
		ļ				AAPALTNPTAPVLCTAPNNTAQKERVPSOMR   ORPAGVRISSRTPDLTCAVSTHSTVPGVRISSC
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						STVPGVRISSCTPDLTCAVSTH
				1		TPDLTCAVSIHATVPGVRISSCTPDLTCAVSIH
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	1					TPDLTCAVSIHSTVPGVRISSCTPDLTCAVSIH
	İ			İ		ATVPGVRISSCTPDLTCAVSTHSTVPGVRISSR
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			l			TPDLTCAVSIHATVPGVRISSRTPDLTCAVSIH
1	}		ľ	1	Ì	ATVPGVRISSCTPDLTCAVSTHSTVPGVRISSR
						TPDLTCAVSIHATVPGVRISSCTPDLTCAVSTH
						STVPGVRISSRTPDLTCAVSIHATVPGVHISSC
ľ	ì	l	ľ	1	Ì	TPDLTCAVSTHSTVPGVRISSRTPDLTCAVSIH
			1			STVPGVCISSRTPDLTCAVSIHSTVPSVHISSCT
						PDLTCAVSIHSTVPGVRISSRTPDLTCAVSTHS
ì		ĺ	ľ	1	Ì	TVPGVHISSCTTDLTCAVSIHATVPGVHISSCT
ļ						PDLTCAVSTHTTVPGVRISSRTPDLTCAVSIHS
						TVPGVRISSCTPDLTCAVSTHSTVPGVRISSRT
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1						TVPGVHISSCTPDLTCAVSIHATVPGVRISSRT
			J	}		PDLTCAVSIHATVPGVHISSCTPDLTCAVSTHS
	İ					TVPGVRISSRTPDLTCAVSIHSTVPGVHISSCT
			ĺ			PDLTCAVSTHSTVPGVHISSCTPDLTCAVSTH
ļ	]	]	ļ	1		STVPGVHISSRTPDLTCAVSIHATVPSVHISSC
						TPDLTCAVSIHSTVPGLLTSVSQTSTG
216	1566	A	2477	1	414	FRTKSYRKGSYRCIVSEWIAEQGNWQEIQEK
l		1	ł			AVEVATVVIQPTVLRAAVPKNVSVAEGKELD
			}			LTCNITTDRADDVRPEVTWSFSRMPDSTLPGS
	Į.					RVLARLDRDFLVHSSPHVALSHVDARSYHLL
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HRAQIMAR YPEGYRTLPRNSKTRPESICSVTP STHDKTLGPGAEEKRRSMRDDTMWQLYEW QQRQFYNKQSTLPRHSTLSSPKTMVNISDQT MHSIPTSPSHGSIAA YQGYSPQRTYRSEVSSPI QRGDVTIDRRHRAHHPKVK  228 1578 A 2583 3 330 LPFLGLGSVLPQGMVMASPEMNPTICSVFEA HIVLLFHATTFRRGFQVTVLVGNVRQTAVVE KIHAKVRGTWPFISPEVRKEGGLPQTGRELLD PTMGIKPHLWWVAA  229 1579 A 2589 1 448 DDKNAQGIKRHVKPTSGNAFTICKYPCGKSR ECVAPNICKCKPGYIGSNCQTALCDPDCKNH	1	1	ļ	)			LEQWIKIQKGRGHEEETRGVISYQTLPRNMPS
STHDKTLGPGAEEKRRSMRDDTMWQLYEW QQRQFYNKQSTLPRHSTLSSPKTMVNISDQT MHSIPTSPSHGSIAAYQGYSPQRTYRSEVSSPI QRGDVTIDRRHRAHHPKVK  228 1578 A 2583 3 330 LPFLGLGSVLPQGMVMASPEMNPTICSVFEA HIVLLFHATTFRRGFQVTVLVGNVRQTAVVE KIHAKVRGTWPFISPEVRKEGGLPQTGRELLD PTMGIKPHLWWVAA  229 1579 A 2589 1 448 DDKNAQGIKRHVKPTSGNAFTICKYPCGKSR ECVAPNICKCKPGYIGSNCQTALCDPDCKNH			1	[			
QQRQFYNKQSTLPRHSTLSSPKTMVNISDQT MHSIPTSPSHGSIAAYQGYSPQRTYRSEVSSPI QRGDVTIDRRHRAHHPKVK  228 1578 A 2583 3 330 LPFLGLGSVLPQGMVMASPEMNPTICSVFEA HIVLLFHATTFRRGFQVTVLVGNVRQTAVVE KIHAKVRGTWPFISPEVRKEGGLPQTGRELLD PTMGIKPHLWWVAA  229 1579 A 2589 1 448 DDKNAQGIKRHVKPTSGNAFTICKYPCGKSR ECVAPNICKCKPGYIGSNCQTALCDPDCKNH				]			
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228 1578 A 2583 3 330 LPFLGLGSVLPQGMVMASPEMNPTICSVFEA HIVLLFHATTFRRGFQVTVLVGNVRQTAVVE KIHAKVRGTWPFISPEVRKEGGLPQTGRELLD PTMGIKPHLWWVAA  229 1579 A 2589 1 448 DDKNAQGIKRHVKPTSGNAFTICKYPCGKSR ECVAPNICKCKPGYIGSNCQTALCDPDCKNH			1		,		
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KIHAKVRGTWPFISPEVRKEGGLPQTGRELLD   PTMGIKPHLWWVAA	220	12/0	ΙΛ.	2303	ا		
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229 1579 A 2589 1 448 DDKNAQGIKRHVKPTSGNAFTICKYPCGKSR ECVAPNICKCKPGYIGSNCQTALCDPDCKNH	ĺ	Ì	ĺ		Ī I		,
ECVAPNICKCKPGYIGSNCQTALCDPDCKNH		1.550	<b> </b>	0.700			
	229	1579	A	2589	1	448	
GKCIKPNICQCLPGHGGATCDEEHCNPPCQH	1	}	l				
	L		L				GKCIKPNICQCLPGHGGATCDEEHCNPPCQH

SEQ ID NO: of	SEQ ID NO: of	Met ·	SEQ ID NO:	Predicted beginning	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide	nou	in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-	ļ	USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence			914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
1				amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
		}		residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
				peptide		/=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion
						GGTCLAGNLCTCPYGFVGPRCETMVCNRHC
230	1580	A	2593	2	138	ENGGQCLTPDICQCKPGWYGPTCSTA  AVTFSVVFAYVADITQEHERSMAYGLVCMFI
230	1300	A	2393		130	LYLLYLLRNAFFLR
231	1581	A	2595	185	2	SGPYTDFTPWPTEEQKLLEQALKTYPVNPPER
221	1501	1.	2373	102	-	WEKIAEAVPGRTKKACIKRYKVADLRISK
232	1582	A	2596	1	391	STVTGQPRRLLDTAGHQQPFLELKIRANEPGA
				_		GRARRTPTCEPATPLCCRRDHYVNFQELGW
						RDWILLPEGYQLNYCSGQCPTHLAGSPGIAAS
	ĺ			1		FHSAVFSLLKANNPWPGRTSWCVPTARRPLS
						LLYL
233	1583	A	2601	184	403	LLFSDEIIMAAPLRIADVTSGLIGGEDGRVYV
						YNGKETTLGDMTGKCKSWITPCPEEKVNVLQ
						NSIPYWERIT
234	1584	A	2614	178	335	PLTLCLPENNKPPQADAVPDKELTLPVDSTTL
025	1505		2616		806	DGSKSSDDQKIISYLWEKTQ
235	1585	A	2616	2	896	DVLEVYGTGVASTRHEMGTLDKHKELEDLV
						AKFLNVEAAMVFGMGFATNSMNIPALVGKG
1						CLILRDEVNHTSLVLGARLLGATIGIFKHNYA   QSLEKLLRDAVIYGQPRTRRAWKKILILVEGV
						YSMEGSIVHLPQIIALKKKYKAYLYIDEAHSI
1						GAVGPTGRGVTEFFGLDPHEVDVLMGTFTKS
						FGASGGYIAGRKARILSPPACLVPNTGSHSLH
	!					RLTRDLQMNEAMVALVTDRLQGWNSGEGN
						WDRADKFGDLVDYLRVHSHSAVYASSMSPPI
						AEQIIRSLKLIMGLDGTTQ
236	1586	A	2621	1	392	NTSSFPAQPSSPARPSLPHLSQHPSNPLLPLAS
						ADHPQCGRFLPLHEPEPLCPSPSLSYPTLVSS
						WSSPFSSHHGCPPGLYPFPTSPKTIQPPGLAQL
						KMLCIPPGRQQLRGAQSMPGHGALSPLLLPP
237	1587	A	2628	398	1	A DLVCKISGFGRGPRDRSEAVYTTMSGRSPAL
237	1301	A	2020	390	1	WAAPETLQFGHFSSASDVWSFGIIMWEVMAF
1						GERPYWDMSGQDVIKAVEDGFRLPPPRNCPN
1						LMHRLMLDCWQKDPGERPRFSQIHSILSKMV
1						QDPEPPNV
238	1588	A	2631	1	1104	WSPCSLTCGVGLQTRDVFCSHLLSREMNETV
						ILADELCRQPKPSTVQACNRFNCPPAWYPAQ
1 1						WQPCSRTCGGGVQKREVLCKQRMADGSFLE
		'			'	LPETFCSASKPACQQACKKDDCPSEWLLSDW
				!		TECSTSCGEGTQTRSAICRKMLKTGLSTVVNS
						TLCPPLPFSSSIRPCMLATCARPGRPSTKHSPHI
						AAARKVYIQTRRQRKLHFVGGGFAYLLPKTA
						VVLRCPARRVRKPLITWEKDGQHLISSTHVT
						VAPFGYLKIHRLKPSDAGVYTCSAGPAREHF VIKLIGGNRKLVARPLSPRSEEEVLAGRKGGP
						KEALQTHKHQNGIFSNGSKAEKRGLAANPGS
						RYDDLVSRLLEQGAPCSSSKKKN
239	1589	A	2636	1	678	MKPDNILLDEHGHVHITDFNIAAMLPRETQIT
						TMAGTKPYMAPEMFSSRKGAGYSFAVDWW
						SLGVTAYELLRGRRPYHIRSSTSSKEIVHTFET
						TVVTYPSAWSQEMVSLLKKLLEPNPDQRFSQ
						LSDVQNFPYMNDINWDAVFQKRLIPGFIPNK
] ]						GRLNCDPTFELEEMILESKPLHKKKKRLAKK
						EKDMRKCDSSQTCLLQEHLDSVQKEFIINRE
240	1500		0(00	200		KVNRDCI
240	1590	Α	2639	389	3	ELLDPTTPMRTKCIELLYAALTSSSTDQPKAD
					}	LWQNFAREIEEHVFTLYSKNIKKYKTCIRSKV
<u> </u>						ANLKNPRNSHLQQNLLSGTTSPREFAEMTVM

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  EMANKELKQLRASYTESCIQEHYLPQVIDGTL Y  IRLTILRCVFMRLATICVLVFTLGSKITSCDDD
			20.0	3,2		TCDLCGYNQKLYPCWETQVGQEMYKLMIFD FIIILAVTLFVDFPRKLLVTYCSSCKLIQCWGQ QEFAIPDNVLGIVYGQTICWIGAFFSPLLPAM Y
242	1592	A	2642	405	1	YFKNTTLLLVGVICVAAAVEKWNLHKRIALR MVLMAGAKPGMLLLCFMCCTTLLSMWLSNT STTAMVMPIVEAVLQELVSAEDEQLVAGNSN TEEAEPISLDVKNSQPSVELIFVNEDILDFLMK SPLMISQACI
243	1593	A	2646	412	2	CLAMIKGIQSSGKIIYFSSLFPYVVLICFLIRAF LLNGSIDGIRHMFTPKLEIMLEPKVWREAATQ VFFALGLGFGGVIAFSSYNKRDNNCHFDAVL VSFINFFTSVLATLVVFAVLGFKANVINEKCIT QNSETV
244	1594	A	2650	1	1271	MTTTLIGLLKTARLLRLVRVARKLDRYSEYG AAVLMLLMCIFALIAHWLACIWYAIGNVERP YLTDKIGWLDSLGQQIGKRYNDSDSSSGPSIK DKYVTALYFTFSSLTSVGFGNVSPNTNSEKIF SICVMLIGSLMYASIFGNVSAIIQRLYSGTARY HMQMLRVKEFIRFHQIPNPLRQRLEEYFQHA WTYTNGIDMNMVTNGTCSSCTSDDGHFILVS NHHQGGLIYSWNDAASMQRPFNHIKSSLLGS TSDSNLNKYSTINKIPQLTLNFSEVKTEKKNSS PPSSDKTIIAPKVKDRTHNVTEKVTQVLSLGA DVLPEYKLQAPRINKFTILHYSPFKAVWDWLI LLLVIYTAIFTPYSAAFLLNDREEQKRRECGY SCSPLNVVDLIVDIMFIIDILINFRTTYVNQNEE VVSDPASV
245	1595	A	2656	385	2	NLTWWPLFRDVSFYIVDLIMLIIFFLDNVIMW WESLLLLTAYFCYVVFMKFNVQVEKWVKQ MINRNKVVKVTAPEAQAKPSAARDKDEPTLP AKPRLQRGGSSASLHNSLMRNSIFQNKIHTLD PHV
246	1596	A	2660	200	506	VLVLQMNYYQMLIIYYVLFFKVNEFLAFEGPI LLDMRIKHLIKTNQLSQATALAKLCSDHPEIG IKGSFKQTYLVCLCTSSPNGKLIEEVSMFSFIS NYFLS
247	1597	A	2678	3	267	DAWVKNDIIFNQTERKQKISENLKHLASVRV VQKNLVFVVGLSQRLADPEVSPLVFFVILIFF VSLSYLEIIFDPAQLCDSSEHIIS
248	1598	A	2687	1	404	DFTTLAAMMRTLFSLFGDVRSDVHRFSVTLF GAAIKSVKNPDKKSIENQVLDSLVPLLLYSQD ENDAVAEESRQVLTICAQFLKWKLPREVYSK DPWHIKPTEAGTICRFFEKKCKGKINILEQTL MYSKNPKL
249	1599	A	2692	1	440	FRRRRRRERDCAAQGARRHCRHLAECKLV SFPIGIYKVLRNVSGQIHLITLANNELKSLTSK FMTTFSQLRELHLEGNFLHRLPSEVSALQHLK AIDLSRNQFQDFPEQLTALPALETINLEENEIV DVPVEKLAAMPALRSINL
250	1600	A	2693	459	21	LLPGSLGVPILHSQPWDPSPQCPHRAPSTPRRL PPLGALSQALTFLSRAAKNHSQDPGKGTKPFP AAPAAPPPRSSLPAPLPMGLKDKGPQPAPPTIF NSPWHPATLPGALGPQLSQAAPSPIPPPCLMG ISSCPDLKLTKSSTP
251	1601	A	2694	2	404	FVFDLKLRVPGFAALLIHGASSVPGPETVRLR

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, I=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  QKRKKKAPDHSSGRKEELVTTHTVDKLETKK
						PVGRVLCGLSGELLHSLLLPRRKTEKRALGSH RKAGFPEHPVAPEPLSNSCQISKEGREQVLSEI GAGDCL
252	1602	A	2697	421	1	PQKSHSGAYQCFATRKAQTAQDFAIIALEDG TPRIVSSFSEKVVNPGEQFSLMCAAKGAPPPT VTWALDDEPIVRDGSHRTNQYTMSDGTTISH MNVTGPQIRDGGVYRCTARNLVGSAEYQARI NVRGPPSIRAMRNIT
253	1603	A	2698	65	401	ACCQWRRTLIPAKSTTVSCTISTPHHPFRGSYS FDDHITDSEALSRSSHVFTSHPRMLKRQPAIEL PLGGEYSSDVPRPLSTQLSSSLLGYFSTLMTG AAFTNNIASSTIIL
254	1604	A	2699	438	301	GQIHSQDDPPFIDQLGFGVAPGFQTFVACQEQ RVRGPWEAGPGVGY
255	1605	A	2700	1	842	LQNREDSSEGIRKKLVEAEELEEKHREAQVS AQHLEVHLKQKEQHYEEKIKVLDNQIKKDLA DKETLENMMQRHEEEAHEKGKILSEQKAMIN AMDSKIRSLEQRIVELSEANKLAANSSLFTQR NMKAQEEMISELRQQKFYLETQAGKLEAQN RKLEEQLEKISHQDHSDKNRLLELETRLREVS LEHEEQKLELKRQLTELQLSLQERESQLTALQ AARAALESQLRQAKTELEETTAEAEEEIQALT VGLGSNIFRLLKASARMSVELALSILAHP
256	1606	A	2701	2	405	FVGGPGADPPVAVMWDPRAARMDLTAYAE LLKESGNQVLKNGNFSLAIRKYDEAIQILLQL YQWGVPPRDLAVLLCNKSNAFFSLGKWNEA FVAAKECLQWDPTYVKGYYRAGYSLLRLHQ PYEAARMFFEGLR
257	1607	A	2702	2	399	FVESASSRPPGCFSGDGRFWLVSEGSRRGWD FNPSFSFLDPRYSVGGDENIGTVTTLANILREF NPSLKGFSVGTGKETSPNAFLNQAVAGGRAE DLPVQARRLVDLMKNDTRIHFQEDWKIITLFI GGNDL
258	1608	A	2709	1	1097	SVGARQGEARDRIRRFFPKGDLEVLQAQVERI MTRKELLTVYSSEDGSEEFETIVLKALVKACG SSEASAYLDELRLAVAWNRVDIAQSELFRGDI QWRSFHLEASLMDALLNDRPEFVRLLISHGLS LGHFLTPMRLAQLYSAAPSNSLIRNLLDQASH SAGTKAPALKGGAAELRPPDVGHVLRMLLG KMCAPRYPSGGAWDPHPGQGFGESMYLLSD KATSPLSLDAGLGQAPWSDLLLWALLLNRA QMAMYFWEMGSNAVSSALGACLLLRVMAR LEPDAEEAARRKDLAFKFEGMGVDLFGECYR SSEVRAARLLLRRCPLWGDATCLQLAMQAD ARAFFAQDGVQSLPTQKWWGDMARR
259	1609	A	2721	1	403	VYLGAGPGLFFSNEGAKEGEKANIPKLMLPR GGFSQREMVTGERSPSPEEEEEEEEGFGERA SCRRGLFRVRLTRVGLAAPSKASRGQEGDAA PKSPVREKSPKFRFPRVSLSPKARSGSGDQEE GGLRVRLP
260	1610	A	2728	1	477	LLGGDLRYHLQQNVHFTEGTVKLYICELALA LEYLQRYHIIHRDIKPDNILLDEHGHVHITDFN IATVVKGAERASSMAGTKPYMAPEVFQVYM DRGPGYSYPVDWWSLGITAYELLRGWRPYEI HSVTPIDEILNMFKVERVHYSSTWCKGMVAL LRK
261	1611	A	2730	3	547	LTITDFILVLYRYYRSPLVQIYEIEQHKIETWR EIYLQGCFKPLVSISPNDSLFEAVYTLIKNRIH

CECID	CEO ID	1 1 / 0+	Tero	D	T 20 11 7 1 1	() ()
SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide	1	in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence	1	J	914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
				amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
	ļ		ļ	residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
1	1	l	1	peptide	Sequence	/=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion
<b> </b>		-		sequence	ļ	
ĺ	1	1		[	[	RLPVLDPVSGNVLHILTHKRLLKFLHIFGSLLP
						RPSFLYRTIQDLGIGTFRDLAVVLETAPILTAL
		1		)		DIFVDRRVSALAVVNECGTHPQDERLGLGW
İ						GLGEPGSEERLFPAAITSR
262	1612	A	2733	3	431	GPEFPGSAKLVFLDLSYNNLTQLGAGAFRSA
		ł				GRLVKLSLANNNLVGVHEDAFETLESLQVLE
						LNDNNLRSLSVAALAALPALRSLRLDGNPWL
J		J	1	ļ	1	
	j			]	J	CDCDFAHLFSWIQENASKLPKGLDEIQCSLPM
	<u> </u>	ļ				ESRRISLRACRRPASRV
263	1613	A	2736	2	343	PARISGVDPPVRKATKGGENCSFEDNKNWQF
	1	l	1		ł	LWGLNGNFNFFKEPWGGRNNHAKGFRTTW
		1	ĺ			ARSSSQNNRTFQNNRNFLRLQRDSQKKGQFA
		İ	1			RLISPLVNLPQSPGGLEFQYQAT
264	1614	A	2738	2	245	RAMLKCLREGOPPPSYNWTRLDGPLPSGVRV
20-1	1014	1	2/30	-	273	
	ļ	1	1		}	DGDTLGFPPLTTEHSGIYVRHDTNEFSSRDSH
<u> </u>	1.7.	<u> </u>	<del> </del>			DTVDVLDPPEDSGKQVDL
265	1615	Α	2752	2	388	AAGDAPLRSLEQANRTRFPFFSDVKGDHRLV
	ľ	1	1		(	LAAVETTVLVLIFAVSLLGNVCALVLVARRR
İ						RRGATACLVLNLFCADLLFISAIPLVLAVRWT
	j	J	] .		J	EAWLLGPVACHLLFYVMTLSGSVTILTLAAV
						SLER
266	1616	A	2755	192	1	AFREVGGYWGLLCEHLYAIPSKTSEGNWTAK
200	1010	A	2/33	192	} 1	
						LQGYLPLQDAFHIFQDPLTGDLPWPELILGLP
						V
267	1617	A	2760	434	714	ASRLEKQNSTPESDYDNTPNDMEPDGMGYM
		ĺ	[			HRTSVPGEGLPRARDLAGLGQQKQFTTHTPF
			1			LYFQTHKGLKDSSIRSEVTCLGISQCWRKGFF
268	1618	A	2762	1	405	IACTFCGQDEWSPERSTRCFRRSRFLAWGEP
				_		AVLLLLLLSLALGLVLAALGLFVHHRDSPL
						VQASGGPLACFGLVCLGLVCLSVLLFPGOPSP
	Ĭ	i	1			
			1			ARCLAQQPLSHLPLTGCLSTLFLQAAEIFVESE
						LPLSWAE
269	1619	A	2772	3	243	TRPAEKIQYLVLFFVMSHPSQAYDKLSLSDHL
						LIAVLNLLRREVSEHGRHLQQYFNLFVMYAN
	1					LSKNLSFSEFCFDVSY
270	1620	Α	2789	1	486	ELQSQQACTHTKETEQLRSQLQTLKQQHQQA
		]	] ]	_	1.55	VEQIAKAEETHSSLSQELQARLQTVTREKEEL
	1	1	1			LQLSIERGKVLQNKQAEICQLEEKLEIANEDR
	1					
١ .	ł	1	1 1		}	KHALERFEQEAVAVDSNLRVRELQRKVDGIQ
	1	1				KAYDELRLQSEAFKKHSLDLLSKERELNGKL
		<u> </u>		-*		RHLSP
271	1621	A	2795	1	568	KEKRVTVQLPTESIQKNQEDKLKMVPRKQRE
			[			FSGSDRGKLPGSEEKNQGPSMIGRKEERLITE
						RKHEHLKNKSAPKVVKQKVIDAHLDSQTQN
		1	1			FQQTQIQTAESKAEHKKLPQPYNSLQEEKCLE
						VKGIQEKQVFSNTKDSKQEITQNKSFFSSVKE
				İ		
270	1.622		0505		500	SQRDDGKGALNIVEFLRKREELHQILSTVKQP
272	1622	A	2797	8	523	KCMQGKYAGAMESEPCVCTEADFDCDYGYE
	J		] ]			RHSNGQCLPAFWFNPSSLSKDCSLGQSYLNST
						GYRKVVSNNCTDGVREQYTAKPQKCPGKAP
						RGLRIVTADGKLTAEQGHNVTLMVQLEEGD
			1		l	VQRTLIQVDFGDGIAVSYVNLSSMEDGIXHV
273	1623	_	2001	70	205	YQNXGIXRXTVQVDNSLGS
213	1023	A	2801	72	395	HPSRSNVGPRQLTVWNTSNLSHDNRRKYIFS
			, !		1	DEEGQNQLGIRIHQDIPLPPRRRELPALRTTNG
			1			KADSLNVSRNSVMQELSELEKQIQVIRQELQL
			1 1			AVSRKTELEEYH
274	1624	A	2805	168	320	ILWLYFETGTWVYPVFAKLSLLGLAALFSLRE
			[			IFIARNGVVGETLTHCKRV
		L				

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275	1625	A	2812	208	321	GSLATCQLSEPLLWFILRVLDTSDALKAFHD MGKIIFQ
276	1626	A	2813	41	266	AGRSLHGAGDRAWVGISPTDWSPKVVELCK KYQQQTVVAIDLAGDETIPGSSLLPGHVQAY OVGPVRRNGEAGPG
277	1627	A	2817	3	410	VLQERLDNFQRKCIQLASSTEGKVDKLLMRN LFISYLHTPKHKQHEVLQAMGSILGITGEEME PLFQEEHGTATRWMTGWLEGGSKSVPKTPL GLNQQPALNGSFSELFVKFLKTESLSSTLPTX LPPHNSPGKIK
278	1628	A	2821	238	457	GLSGPSCSCPHSPLPTIISRAQLETALKWRNYE VKLRLLLHLEELQMEHDIRHYDLESVPMTWD PVDONPRLV
279	1629	A	2822	342	1	PLIPANLPAHSNPLQPLPSLPHPFLPATHKFPT TPPTFSSVPPPLPSLSSILHHSPLHSELNPHLQS CRLPSRPSVSRELPPQSGPASSVPLAPTPLPDS VPSQRHPTXPPPAS
280	1630	A	2825	307	77	PSMVWSYHWGVKQKRLALCVFSFEEGGRRK CGQYWPLEKDSRIRFGFLTVTNLTGAVGEPG VAFQCDGQRRREPTC
281	1631	A	2827	81	381	KMGTAVWVPKEKEKRDKASQEGGDVLGAR QDCTPSLKSLVATGNLLDLEETAKAPLSTVSA NTTNMDEVPRPQALSGSSVVWVSGCVASRS VILSLTSG
282	1632	A	2830	471	160	KLPXDKYELEPSPLTQYILERKSPHTCWQVFV TSSGKYNELGYPFGYLKASTTLTCVNLFVMP YNYPVLLPLLDDLFKVHKLKPNLKWRQAFDS YLKTLPPYYL
283	1633	A	2835	462	148	VSPALSLTPTIFSYSPSPGLSPFTSSSCFSFNPEE MKHYLHSQACSVFNYHLSPRTFPRYPGLMVP PLQCQMHPEESTQFSIKLQPPPVGRKNRERVE SSEESAP
284	1634	A	2836	2	384	KTLPRTLLDILADGTILKVGVGCSEDASKLLQ DYGLVVRGCLDLRYLAMRQRNNLLCNGLSL KSLAETVLNFPLDKSLLLRCSNWDAETLTED QVIYAARDAQISVALFLHLLGYPFSRNSPGEK KR
285	1635	A	2843	20	271	PIRPYYSYSGLDRDCSWLPLAKAWLPDVMIL VCDRVSEDGINRQQAQEWCIKHGFELVELSP EELPEEDGKCLCVRRKYGTYI
286	1636	A	2845	197	278	TAEDVLTVAYEHGVNLFDTAEVYAAGK
287	1637	A	2851	2	427	FVAEVRREWAKYMEVHEKASFTNSELHRAM NLHVGNLRLLSGPLDQVRAALPTPALSPKDK AVLQNLKRILAKVQEMRDQRVSLEQQLRELI QKDDITGSLVTTDHSQMKKLFEEQLKKYDQL KVYLEQNLAAQDRVLCALT
288	1638	A	2859	2	469	FVNLGILTCIECSGIHREMGAHISRIQSLELDK LGTSELLPAKNVGNNSFNDIMEANLPSPSPKP TPSSDMTVRKEYITAKYVDHRFSRKTCSTSSA KLNELLEAIKSRDLLALIQVYAEGVELMEPLL EPGQELAETALHLAVRTADQTSLHLVE
289	1639	A	2861	2	454	FVASGGPATARMSDSQFFCVAEERSGHCAVV DGNFLYVWGGYVSIEDNEVYLPNDEIWTYDI DSGLWRMHLMEGELPASMSGSCGACINGKL YIFGGYDDKGYSNRLYFVNLRTRDETYIWEK ITDFEGQPPTPRDKLSCWVYKDRLIYFG
290	1640	A	2868	1	378	FRQGQLYKVFLHGSQGQVYHSQQVGPPGSAI SPDLLLDSSGSHLYVLTAHQVDRIPVAACPQF PDCASCLQAQDPLCGWCVLQGRCTRKGQCG

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		ĺ	l			RAGQLNQWLWSYEEDSHCLHIQSLLPGHHPR QE
291	1641	A	2870	1	385	FRYMPNNRQQLLRKRHIGNDIVTIVFQEPGAL PFTPKSIRSHFQHVFVIVKVHNPCTENVCYSV GVSRSKDVPPFGPPIPKGVTFPKSAVFRDFLL AKVINAENAAHKSEKFRAMATRTRQEYLKD LA
292	1642	A	2877	3	188	RPTRPPPATTQSPESTMDTSLKKEKSAILDLYI PPPPAVPYSPRYVAVHCHGMLVSCWCHL
293	1643	A	2878	1	427	REKEEEVEEEEDKVVKETEKEAEQEKEEDSL GAGTHPDAAIPSGERTCGSEGSRSVLDLVNYF LSPEKLTAENRYYCESCASLQDAEKVVELSQ GPCYLILTLLRFSFDLRTMRRRKILDDVSIPLL LRLPLAGGRGQAYDL
294	1644	A	2879	109	245	QLCCFCFRQTTLIVYILSFIGMVIFTFTLDLRYI IIVFVTGGVLG
295	1645	A	2880	3	320	LASSQHGILNNLSLLFSICKTCIRTMDHHCPRA NNCVGEQNHRFFCALHCKSKHFCIEFTLNTNF FNCFLPGAEKSTIDAPFSLQPFLQDSKYNTALS LSESISQ
296	1646	A	2892	209	363	SQYSHSLDYHLLQVTKNPFTLGDSSNPGQTE RLQEFSQKMDQVRGHWPVST
297	1647	A	2893	8	424	SPXTLXLDTFILLGIQDNILVLILATPPFMAGG KLYSTMGRFLRDRKNPACREMAVVLLANLA QGDSLAARAIAVQKGSIGHLLGFLEDSLAAT QIQQSQASLLHMHNPPFEPTSVDMMRRACRA LLALAKVDDNHSEF
298	1648	A	2894	310	445	FWIYFPSFFMTGYLPLGFEFAVEITYPESEGTS SGLLNASAQVNL
299	1649	A	2898	1	492	KIKAKNLTNYDLCSIFLGTSTLLVWVGVIRYL GYFQAYNVLILTMQASLPKVLRFCACAGMIY LGYTFCGWIVLGPYHDKFENLNTVAECLFSL VNGDDMFATFAQIQQKSILVWLFSRLYLYSFI SLFIYMILSLFIALITDSYDTIKKFQQNGFPETD LQEF
300	1650	A	2901	1	445	PVWWNSLNGASEVTFSVHVKDGGSFPKTDST TVTVRFVNKADFPKVRAKEQTFMFPENQPVS SLVTTITGSSLRGEPMSYYIASGNLGNTFQIDQ LTGQVSISQPLDFEKIQKYVVWIEARDGGVPP FSSYEKLDITVLDVNDNAPIF
301	1651	A	2902	162	433	THFICLPLGYCFPLLDKDLQLPSGFNCNFDFLE EPCGWMYDHAKWLRTTWASSSSPNDRTFPG KPAVSEDMKELRPACSTYFNPRFPYKL
302	1652	A	2909	2	412	GPQMLCKKIYFIWVTRSQCQFEWLADIMQEV EENDHQDLVSVHIYVTQLAEKFDLRTTMLYI CERHFQKVLNRSLFTGLRSITHFGRPPFEPFFN SLQEVHPQVRKIGVFSCGPPGMTKNVEKACQ LVNRQDRAHFM
303	1653	A	2914	291	453	KLNRWLCFFYSWSFGILLYEMVTLGAPPYPE VPPTSILEHLQRRKIMKRPSSCS
304	1654	A	2926	179	354	PGVPSQALRKAESLKKCLSVMEAKVKAQTAP NKDVQREIADLGEVGAASLPPSSGPGA
305	1655	A	2938	135	438	GMGYLHAKGILHKDLKSKNVFYDNGKVVIT DFGLFSISGVLQAGRREDKLRIQNGWLCHLA PEIIRQLSPDTEEDKLPFSKHSDVFALGTIWYE LHAREWP
306	1656	A	2944	2	329	VRWNSCVNCSCAFGNGASLSTSLGESSGCLW EIGKWLSCSLLSFPSPLAVLIITFCIVTVLGREA LTKGALWAVFLLAGSALLCAEVTGVIWRQPE

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307	1657	A	2950	2	411	SKTKLSFKVSSSA  NYLCIAKNSAGSAMGKTRLVVQVPPVIENGL PDLSTTEGSHAFLPCKARGSPEPNITWDKDGQ PVSGAEGKFTIQPSGELLVKNLEGQDAGTYT CTAENAVGRARRRVHLTILVLPVFTTLPGDRS LRLGDRLWLR
308	1658	A	2951	1	407	PTRPPRVRFDNEFDAESQRKRTTSVSKMERM DSSLPEEEEDEDKEAINGSGNAENRERHSESS DWMKTVPSYNQTNSSMDFRNYMMRDETLEP LPKNWEMAYTDTGMIYFIDHNTKTTTWLDP RLCKKAKAPEDC
309	1659	A	2954	2	179	QDFLTLTLTEPTGLLYVGAREALFAFSMEALE LQGAVRGGAVGGSRACQRARPRGAVLG
310	1660	A	2959	1	419	QDMMERAIIDTFVGHDVVEPGSYVQMFPYPC YTRDDFLFVIEHMMPLCMVISWVYSVAMTIQ HIVAEKEHRLKEVMKTMGLNNAVHWVAWFI TGFVQLSISVTALTAILKYGQVLMHSHVVIIW LFLAVYAVATIMFCF
311	1661	A	2963	3	465	MKPQMPGLGAPNGYGPGRGRAGVPGGPERR PWVPHLLPFSSPGYLGVMKAQKPGAGEGMK PQKPGLRGTLKPQKSGHGHENGPWPGPCNA RVAPMLLPRLPTPGVPSDKEGGWGLKSQPPS AVQNGKLPGHQPPNGYGPGAEPGFNGGLEPQ KI
312	1662	A	2967	3	405	WLAQEWSPCTVTCGQGLRYRVVLCIDHRGM HTGGCSPKTKPHIKEECIVPTPCYKPKEKLPV EAKLPWFKQAQELEEGAAVSEEPSFIPEAWS ACTVTCGVGTQVRIVRCQVLLSFSQSVADLPI DECEGPKPA
313	1663	A	2969	2	430	VVADNCRQGYLDALRFLERRGLTKEPVLWT LVSKEPPAPADGNWDAGCDQRRKGGLSLNW KVPHVQVKDVPNFEQLSPELEAALKKACTRD PSRWARFWHSGPGQVLTYLLLPCTLPFEYIYF RSRRLVVWLPDVPADLWWMQ
314	1664	A	2971	422	33	LDXSHNALQRLRPGWLAPLFQLRALHLDHNE LDALGRGVFVNASGLRLLDLSSNTLRALGRH DLDGLGALEKLLLFNNRLVHLDEHAFHGLRA LSHLYLGCNELASFSFDHLHGLSATHLLTLDL SSNRM
315	1665	A	2973	1	525	ITVSTHASGSPFGLEPQSGWLWVRAALDREA QELYILKVMAVSGSKAELGQQTGTATVRVSI LNQNEHSPRLSEDPTFLAVAENQPPGTSVGRV FATDRDSGPNGRLTYSLQQLSEDSKAFRIHPQ TGEVTTLQTLDREQQSSYQLLVQVQDGGSPP RSTTGTVHVAVLDLNDNT
316	1666	Α	2978	2	400	ELVVELVSAGKSGPERNTYEVQVVTGNVPKA GTDANVYLTIYGEEYGDTGERPLKKSDKSNK FEQGQTDTFTIYAIDLGALTKIRIRHDNTGNR AGWFLDRIDITDMNNEITYYFPCQRWLAVEE DDGQLSRE
317	1667	A	2981	3	440	VLNCQGRPTRPVRINGDGQEVLYLAESDNVR LGCPYVLDPDDYGPNGLDIEWMQVNSNPAH HRENVFLSYQDKRINHGSLPHLQHRVRFAAS DPSQYDASINLMNLQVSDTATYECRVKKTTM ATRKVIVTVQARPAVPMCWTEGQ
318	1668	A	2995	119	414	LPEKEFPIIRKSSSLKVTKCLFTEQPKPIIILRFA ENYDARI.LRIDIANTLREQVQELFNKTYGKQ RRTPGEGHVAAVDREVAGFPVPAEGISGETIH
319	1669	Ā	2999	2	332	GFFAYTYGRLVVVEDLHSGAQQHWSGHSAEI

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200	1670		2000		000	VSGGLCQHLIFPHSTTVLALAFSPDDRLLVTL GDHDGRTLALWGTGHL
320	1670	A	3000	693	322	IDESTGLIITVNYLDYETKTSYMMNVSATDQA PPFNQGFCSVYITLLNELDEAVQFSNASYEAA ILENLALGTEIVRVQAYSIDNLNQITYRFDAY TSTQAKALFKIDAITVRGWGQGAPFFPI
321	1671	A	3001	6	383	RIPRGKACXTVLGRSTGELEGFASSRLPPQPC GWGQSSDLLSRIDLDELMKKDEPPLDFPDTLE GFEYAFNEKGQLRHIKTGEPFVFNYREHLHR WNQKRYEALGEIITKYVYELLEKDCNSKKVS
322	1672	A	3007	192	447	ERVRNSLFPGRGDSQCACCPSSPVWVFLETGF LFPWLFLQVEVIKKAYMQGEVEFEDGENGK DGAASPRNVGHNIYILAHQLARH
323	1673	A	3019	18	245	KELLFYHLIVNNINFFNTRYAKIHIPIIASVSEH QPTTWVSFFFDLHILVCTFPAGLWFCIKNIND ERVFGKRGF
324	1674	A	3020	523	797 .	LCYFSARYHQRKIFGILYIFTLSAINRKEPNLFI YLFIFFEMESHSVTHAGVQRHNLNSLQPLPPG FKRFSCLCFLSSWNYRGAPPGPANF
325	1675	A	3022	2	156	NDFLPLYFGWVLTKKSSETLRKAGQVFLEEL GNHKAFKKELRQCRWQVGAL
326	1676	A	3023	38	172	KMVRGSKKLISFFPGGPYGILAGRDPSKGLAT FCLNKEALKDEFE
327	1677	A	3027	1	385	LTLEFLLLPAASELAHGKRLACCIVDHKLPEC GFYGLYDKILLFKHDPTSANLLQLVRSSGDIQ EGDLVEVVLSASATFEDFQIRPHALTVHSYRA PAFCDHCGEMLFGLVRQGLKCDGCGLNYHK RC
328	1678	A	3030	13	569	ITRPTISCQRPGPGLAAGMLPYTVNFKVSART LTGALNAHNKAAVDWGWQGLIAYGCHSLV VVIDSITAQTLQVLEKHKADVVKVKWAREN YHHNIGSPYCLRLASADVNGKIIVWDVAAGV AQCEIQEHAKPIQDVQWLWNQDASRDLLLAI HPPNYIVLWNADTGTKLWKKSYADNILSFSF D
329	1679	A	3038	90	744	SVNLPPSLWPWEEAMDSTKSEPLKGSPEAED GNIEYKKLVNPSQYRFEHLVTQMKWRLQEG RGEAVYQIGVEDNGLLVGLAEEEMRASLKTL HRMAEKVGADITVLREREVDYDSDMPRKITE VLVRKVPDNQQFLDLRVAVLGNVDSGKSTL LGVLTQGELDNGRGRARLNLFRHLHEIQSGR TSSISFEILGFNSKGEVHGINGTQWGQTLRMG W
330	1680	A	3040	3	397	LCSTLLLLTIPSWVLSQITLKESGPTLMKPTET LTLTCTFSGFSLNTSGVGVAWIRQPPGKALE WLALIYWDDDKRYSPSLNDRLTIAKDTSRNQ VVLTMTNMGPVDTATYYCAQFARGARGSN WFDPWGQ
331	1681	A	3043	3	1509	AGIRHEAPPTTSNRHRRQIDRGVTHLNISGLK MPRGIAIDWVAGNVYWTDSGRDVIEVAQMK GENRKTLISGMIDEPHAIVVDPLRGTMYWSD WGNHPKIETAAMDGTLRETLVQDNIQWPTG LAVDYHNERLYWADAKLSVIGSIRLNGTDPI VAADSKRGLSHPFSIDVFEDYIYGVTYINNRV FKIHKFGHSPLVNLTGGLSHASDVVLYHQHK QPEVTNPCDRKKCEWLCLLSPSGPVCTCPNG KRLDNGTCVPVPSPTPPPDAPRPGTCNLQCFN GGSCFLNARRQPKCRCQPRYTGDKCELDQC

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						FLGDRCQYRQCSGYCENFGTCQMAADGSRQ CRCTAYFEGSRCEVNKCSRCLEGACVVNKQS GDVTCNCTDGRVAPSCLTCVGHCSNGGSCT MNSKMMPECQCPPHMTGPRCEEHVFSQQQP GHIASILIP
332	1682	A	3045	3	952	TTTISNFHTQVNRTYCCGTYRAGPMRQISLVG AVDEEVGDYFPEFLDMLEESPFLKMTLPWGT LSSLRLQCRSQSDDGPIMWVRPGEQMIPTAD MPKSPFKRRSMNEIKNLQYLPRTSEPREVLF EDRTRAHADHVGQGFDWQSTAAVGVLKAV QFGEWSDQPRITKDVICFHAEDFTDVVQRLQ LDLHEPPVSQCVQWVDEAKLNQMRREGIRY ARIQLCDNDIYFIPRNVIHQFKTVSAVCSLAW HIRLKQYHPVVEATQNTESNSNMDCGLTGKR ELEVDSQCVRIKTESEEACTEIQLLTTASSSFP PASE
333	1683	A	3046	497	167	SACSTGPELPGRATRSLTRPANQKGCDGDRL YYDGCAMIAMNGSVFAQGSQFSLDDVEVLT ATLDLEDVRSYRAEISSRNLAVSAPVDTCVG CSSKTWKVAPFVRAWWRP
334	1684	A	3053	37	276	VITDLEEQLNQLTEDNAELNNQNFYLSKQLD EASGANDEIVQLRSEVDHLRREITEREMQLTS QKQVRRVNKVVRSLEDF
335	1685	A	3054	2	846	WDAWGDWSDCSRTCGGGASYSLRRCLTGR NCEGQNIRYKTCSNHDCPPDAEDFRAQQCSA YNDVQYQGHYYEWLPRYNDPAAPCALKCH AQGQNLVVELAPKVLDGTRCNTDSLDMCISG ICQAVGCDRQLGSNAKEDNCGVCAGDGSTC RLVRGQSKSHVSPEKREENVIAVPLGSRSVRI TVKGPAHLFIESKTLQGSKGEHSFNSPGVFVV ENTTVEFQRGSERQTFKIPGPLMADFIFKTRY TAAKDSVVQFFFYQPISHQWRQTDFFPCTVT CGGG
336	1686	A	3058	54	347	VVGKQEAGAHSDSCCLLHTPPRLTPAHSRKA LRNSRIVSQKDDVHVCIMCLRAIMNYQVSRG AWDWRLGSPACPHWGLHKLPRLWDPLSLYP VLCWGT
337	1687	A	3059	2	709	ILTSLVELTRFETLTPRFSATVPPCWVEVQQE QQQRRHPQHLHQQHHGDAAQHTRTWKLQT DSNSWDEHVFELVLPKACMVGHVDFKFVLN SNITNIPQIQVTLLKNKAPGLGKVNGLRLCPF LEDHKEDILCGPVWLASGLDLSGHAGMLTLT SPKLVKGMAGGKYRSFLIHVKAVNERGTEEI CNGGMRPVVRLPSLKHQSNKGYSLASLLAK VAAGKEKSSNVKNENTSGTRK
338	1688	A	3060	85	384	KAFYNYHVLELLQMLVTGGVSSQLEQHLDK DKVYGVADSCTSLLSGRNRCKLGLLSLHETIL SDVNPRNTFGQLFCGSLDLFGILCVGLYRIIDE EELNP
339	1689	A	3063	236	362	CFLCLSGDFMVMTIFFNVSRRFGYVAFQNYV PSSVTTMLSWV
340	1690	A	3065	3	1249	DLWQFTPLHEAASKNRVEVCSLLLSYGADPT LLNCHNKSAIDLAPTPQLKERLAYEFKGHSLL QAAREADVTRIKKHLSLEMVNFKHPQTHETA LHCAAASPYPKRKQICELLLRKGANINEKTKE FLTPLHVASEKAHNDVVEVVVKHEAKVNAL DNLGQTSLHRAAYCGHLQTCRLLLSYGCDPN

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0.41	1601		2050			HGADPTKKNRDGNTPLDLVKDGDTDIQDLLR GDAALLDAAKKGCLARVKKLSSPDNVNCRD TQGRHSTPLHLAGK
341	1691	A	3070	1	547	GVLIPSFQNQLFADILAGIESVTSEHNYQTLIA NYNYDRDSEEESVINLLSYNIDGIILSEKYHTI RTVKFLRSATIPVVELMDVQGERLDMEVGFD NRQAAFDMVCTMLEKRVRHKILYLGSKDDT RDEQRYQGYCDAMMLHNLSPLRMNPRAISSI HLRMQLMRDALSANPDLDGVFCTN
342	1692	A	3073	463	3	RINRCRKPSDADILVPGDTISLIGTTSLRIDYNE IDDNRVTAEEVDILLREGEKLAPVMAKTRILR AYSGVRPLVASDDDPSGRNVSRGIVLLDHAE RDGLDGFITITGGKLMTYRLMAEWATDAVC RKLGNTRPCTTADLALPGSQEPAKVP
343	1693	A	3075	250	1	LLIYLAIFAPVAMSALAGVKSVQQVRIRAAQS LGASRAQVLWFVILPGALPEILTGLRIGLGVG WSTLVAAELIAATRGLGFM
344	1694	A	3076	2	138	LYFDAYLQSLQVAAISTFCCLLIGYPLAWAV AHSKPSTRNILLLL
345	1695	A	3078	469	3	LKIRGQRIELGEIDRVMQALPDVEQAVTHAC VINQAAATGGDARQLVGYLVSQSGLPLDTSA LQAQLRETLPPHMVPVVLLQLPQLPLIANGKL DRKALPLPELKAQAPGRAPKAGSETIIAAAFS SLLGCDVQDADADFFALGGHSLLAMKLAT
346	1696	A	3082	404	2	QNITSKDLDVRLDPQTVPIELEQLVLSFNHMI ERIEDVFTRQSNFSADIAHEIRTPITNLITQTEI ALSQSRSQKELEDVLYSNLEELTRMAKMVSD MLFLAQADNNQLIPEKKMLNLAHEVGKVFD QFEALPE
347	1697	A	3084	3	340	NELTFKEAEISKLYTKVHPAYRTLLEKRQALE DEKAKLNGRVTAMPKTQQEIVRLTRDVESGQ QVYMQLLNKEQELKITEASTVGDVRIVDPAIT QPGVLKPKKGLIILGAI
348	1698	A	3086	723	10	TQAMVWQQKACAEDDPQLSGRHWLHAATL YNIAAYPHLKGDDLAEQAQALSNRAYEEAA QRLPGTMRQMEFTVPGGAPITGFLHMPKGDG PFPTVLMCGGLDAMQTDYYSLYERYFAPRGI AMLTIDMPSVGFSSKWKLTQDSSLLHQHVLK ALPNVPWVDHTRVAAFGFRFGANVAVRLAY LESPRLKAVACLGPVVHTLLSGLKCQQVPE MYLDVLASRLGMHDASTKSSTRENH
349	1699	A	3087	2	249	RIRSSDPEITLAGTPLHAAYLIGMTLICAGFSV GFGVAMSQALGPFSLRAGVASSTLGIAQVCG SSLWIWLAAVVGIGAWNM
350	1700	A	3099	3	424	EAPEATPQPSQPGPSSPISLSAEEENAEGEVSR ANTPDSDITEKTEDSSVPETPDNERKASISYFK NQRGIQYIDLSSDSEDVVSPNCSNTVQEKTFN KDTVIIVSEPSEDEESQGLPTMARRNDDISELE DLSGMEDLK
351	1701	A	3108	2	404	IKKNHIIGYQLLHRRALFEKRTRLSDYALIFG MFGIVVMVIETELSWGAYYKAPLYSLALKCL ISLFTIILLGLTIVYHAREIQLFMANYGADDWR SALTYEPIFLILLEALRGVIHATPCRVSLSLWD GLDLP

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352	1702	A	3110	341	2	AQLAEVCPPQTLLTTNTSSISITAIAAEIKNPER VAGLHFFNPAPVMKLVEVVSGLATAAEVVE QLCELTLSWGKQPVRCHSTPGFIVNRVARPY YSEAWRALEEQVAAPEVI
353	1703	A	3111	3	188	HFSLFRIAFAVFLTYMTVGLPLPVIPLFVHHEL GYGNTMVGIAVGIQFLATVLTRGYAGRLA
354	1704	A	3116	367	225	WQLFHLNGTFLNIGETDTESCVNGWVYDRSS FPFSNMTEVRGLVFLS
355	1705	A	3117	101	53	VINLVYLISSPRPELKPVDKESEVVMKFPDGF EKFSPPILQLDEVDFYYDPKHVIFSRLSVSADL ESRICVVGENGAGKSTMLKLLLGDL\APVRGI RHAHRNLKIGYFSQHHVGAAGT*TFSACGNL LGTQVFLGRPEEEY\RHQLGFGMGISGELGHA SSLPACLGGQKEAEVAFCSDGLLPCPNFL\IL\ DEPTN\HLGHGRAIEALGPCLQTISGVGVILVS HE*SALSRLVCRE\LWVC*GRSTSPF
356	1706	A	3121	137	466	RGGRDWGEHNQRLEEHQARAWQGAMDAG AASREHARWQGTGLAPGTRVAVAPTCVQGL PQERSVCRPFFSSRWREGPVWALGAGAHGKP RWSGGVRCVVRGGRWFTPAPH
357	1707	A .	3124	1249	229	MLEAPGPSDGCELSNPSASRVSCAGQMLEVQ PGLYFGGAAAVAEPDHLREAGITAVLTVDSE EPSFKAGPGVEDLWRLFVPALDKPETDLLSH LDRCVAFIGQARAEGRAVLVHCHAGVSRSV AIITAFLMKTDQLPFEKAYEKLQILKPEAKMN EGFEWQLKLYQAMGYEVDTSSAIYKQYRLQ KVTEKYPELQNLPQELFAVDPTIVSQGLKDE VLYKCRKCRRSLFRSSSILDHREGSGPIAFAH KRMTPSSMLTTGRQAQCTSYFIEPVQWMESA LLGVMDGQLLCPKCSAKLGSFNWYGEQCSC GRWITPAFQIHKNRVDEMKILPVLGSQTGKI
358	1708	A	3127	816	139	EVETLGPRTPGP/EAQSPTPGSCPGWQEPSPGP TPPP*LSGPGPQGAPVLGKLLPDPEETPAGKTP LGKHFWWGL\PVTSANFSPGAAA*FGGALSPP GGDL/GHMLLQGPPSPFRLQQQ*QTPPGSHSP PTANREINPGPAAAADTRSCWGHKRSWRGW RGLAPWRLGFGSPGIP*PAPAGIP/GRPTWEGG KGAGGKPSETLTRSPPVWRGKRGSANGFLSW VQILQ
359	1709	A	3132	3	191	HEHLLLLLCVFLVKSQGVNDNEEGFFSARG HRPLDKKREDAPNLRPALAD\ITVCDYRAQIA *AASTPKRAASIAHNAVSCR*AQIA
360	1710	A	3134	1	286	REPPRPALLFF*DRVSLCCPGWNAVVQSQLT AAPTSQVQ/SDSPTFPSSWDYRHVPEYPANFL *RQGFPMLPRLVSNSWAQTVHPPRPPKVLDL QA
361	1711	A		56	1449	PVPAPRVSPSARGAPGRPRLPGVRGPRHS/WA AD*RGSRM/PPRAPAPSPTGP/APGGKKVRGR VPEDPDAYEPRCSAL*V*PTHVTSPQFCDP*N GQIRSYFTVLLRGLNETMLVK/PLCRREP/PEA GPGRQSTPAVTRDHRQHEDPRGAGRQWDAD PRPSAP/PAEVATGSRPGRHMWMRLCLAAQQ APGLPHRTSIRPGWRRLTEPEAWARRHRRW GQRGAVRPPPQGAAPPPSHQGRRTNTDPSAT PRLTVMSRCLAPDLKAPASGPRGWRRGMPQ SS/GALLWTPPPTPRGSHSPRPREAPLRAIHPA GPSK/SRAGASGRLPEVIYGWVTLFTPPEAGT F/LIPSPT*MSPALVIQPPVPPTQMGLRISGLPR QG*PSGAPW*LPGLAQLAFQCHLPHDEVGPP

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						GHSSPREPQVLKKPLWGQTDIAGVGSASLYP DNL
362	1712	A	3136	1270	274	RVGMVLGTREVGDSTPPPSPPLYPFTGNEFVQ HNTWQLSRVYPSDLRTDSSNYNPQELWNAG CQM/V*GGSRDWEEGVEEQQVGNKFSSDGR VGECSRKLLG*EMLSVDITSRYRAPSTYLLNS LKEGLEGLHGESCSSFLLGPSVAMNMQTAGL EMDICDGHFRQNGGCGYVLKPDFLRDIQSSF HPEKPISPFKAQTLLNQVISVQQLPKVDKTKE GSIVDPLVKVQIFGVRLDTARQETNYVENNG FNPYWGQTLCFRVLGPDFPMLRFGKMDYDW KSRNDLLGKTPCPGTCMQQGYRHIHLLSKDG ISLRPASIFVYICIQEGLEGDES
363	1713	С	3139	60	248	MFAGSYGKSMFSFSKKVLNCLPKWRYHFVIA
364	1714	A	3140	57	418	PAMNESPLAPHLHQHLVFSVFQVLTILIGV**  SAFKTLQLPAFSLYFDLGSLKLLILRIHTSIVK NHKVESPRTMSPG*DPQSFLQIPQPRPPQLRV GLTSGLIQHFHSPSSCQFPLLRGPPFPRQPPLGI SGASLCPVLSPPR*PLQPSSL
365	1715	A	3145	122	413	LLPYPSLFVFLRQCHFVTNRLECNGVVSAHCN LHLPGSSDSPASAS*VAGTTGVCHHTRLIF\VF LV*TGFHYVAQAGLELLTA*S\PPQLPKVVGL QA
366	1716	A	3150	247	2	VGEKLHDIRFGNDFDMTPKAQATKEKIDKLN FIKIKKLCIEGYY/NREPQNGRKIFANYVS\DK GLMATIYEELLKLSNKLIQ
367	1717	A	3152	3	2367	QKLKQNQPKRAHVEDGGSRSKQGNEQSKKT PIEKSDFAAATHPRAFYLSKPDETPNAWMSD SGTGLTYWKLEEKDMHHSLPETLEKTFISLSS TDVSPNQVLTLDPTLHMKPKQQISGIQPHGLP NALDDRISFSPDSVLEPSMSSPSDIDSFSQASN VTSQLPGFPKYPSHTKASPVDSWKNQTFQNE SRTSSTFPSVYTITSNDISVNTVDEENTVMVAS ASVSQSQLPGTANSVPECISLTSLEDPVILSKIR QNLKEKHARHIADLRAYYESEINSLKQKLEA KEISGVEDWKITNQILVDRCGQLDSALHEATS RVRTLENKNNLLEIEVNDLRERFSAASSASKI LQERIEEMRTSSKEKDNTIIRLKSRLQDLEEAF ENAYKLSDDKEAQLKQENKMFQDLLGEYES LGKEHRRVKDALNTTENKLLDAYTQISDLKR MISKLEAQVKQVEHENMLSLRHNSRIHVRPS RANTLATSDVSRRKWLIPGAEYSIFTGQPLDT QDSNVDNQLEETCSLGHRSPLEKDSSP/GSSST SLLIKKQRETSDTPIMRALKELDEGKIFKNWG TQTEKEDTSNSLL*/INPRQTETSVNASRSPEK CAQQRGKRLNSASQRSSSLPPSNRKSSTPTKR EIMLTPVTVAYSPKRSPKENLSPGFSHLLSKN ESSPIREKTYSEKATDNHVNHSSCPEPVPNGV KKVSVRTAWEKNKSVSYEQCKPVSVTPQGN DFEYTAKIRTLAETERFFDELTKEKDQIEAAL SRMPSPGGRITLQTRLNQVKCLSLNIL
368	1718	A	3163	2 -	2350	EFKSGGCGAGLVAAGAVLVLYPASRAGERT RVPGSPAPSSLPLHSPGACGTEVDMDPQRSPL LEVKGNIELKRPLIKAPSQLPLSGSRLKRRPDQ MEDGLEPEKKRTRGLGATTKITTSHPRVPSLT TVPQTQGQTTAQKVSKKTGPRCSTAIATGLK NQKPVPAVPVQKSGTSGVPPMAGGKKPSKRP AWDLKGQLCDLNAELKRCRERTQTLDQENQ

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						QLQDQLRDAQQQVKALGTERTTLEGHLAKV QAQAEQGQQELKNLRACVLELEERLSTQEGL VQELQKKQVELQEERRGLMSQLEEKERRLQT SEAALSSSQAEVASLRQETVAQAALLTEREER LHGLEMERRRLHNQLQELKGNIRVFCRVRPV LPGEPTPPPGLLFPSGPGGPSDPPTRLSLSRSD ERRGTLSGAPAPPTRHDFSFDRVFPPGSGQDE VFEEIAMLVQSALDGYPVCIFAYGQTGSGKTF TMEGGPGGDPQLEGLIPRALRHLFSVAQELSG QGWTYSFVASYVEIYNETVRDLLATGTRKGQ GGECEIRRAGPGSEELTVTNARYVPVSCEKEV DALLHLARQNRAVARTAQNERSSRSHSVFQL QISGEHSSRGLQCGAPLSLVDLAGSERLDPGL ALGPGERERLRETQAINSSLSTLGLVIMALSN KESHVPYRNSKLTYLLQNSLGGSAKMLMFV NISPLEENVSESLNSLRFASKVEPSVLFGTAQS NRKWKTDPDLCVCVCVCVCVCVCVCVCVP MSMYRVRGGRVAGGCFIGWRAPCPRAIK
369	1719	Α	3165	365	12	GYTSQGRWIDIERGPLTANTESLIIENNFNALP GYIRKIE*I*IYKKN*INFGGVGLLNIVKISILS/K IYRFDAIPVKILTRFFINLDKLILKFVLKTKIAK NRIKTFYIMRRKKLGDSS
370	1720	Α.	3170	393	42	GASISPSAVIDGVEGLKPMQEQEAQEAGPCLD *HMAPEQWVAPR\RLLFRLIFSVLHALIIAAAA QSSAEEDEDPRN*GQSSEDQAPNQNGLIVIVH RVHVPLGAAATVPVHRSHFPR
371	1721	A	3173	770	510	GNGGCGLSQIPPSHLGAF\$RGSLL\$RG\DPRGP PPHPVIFFVFVVE\QGFTVLARMV\$IS*PCDPP ALA\$Q\$AGITGV\$HLARPQNLYF
372	1722	A	3180	381	76	RVLHHDNVPAHSSPQKREISQEFQLEIRHLP*S PDLAPSGCFLFLNLKNIFK\GTHFSLVDNVKK TVSTWLH/SQNAQFYKDRLNGWYHCLQKCL QHY*AYVEK
373	1723	A	3181	410		RREVAGPEGKGLLLASAHTMLTPPLLLLPLL SALVAAAIDAPKTCSPKQFACRDQITCISKGW RCDGERDCPDGSDEAPEICPQSKAQRCQPNE HNCLGTELCVPMSRLCNGVQDCMDGSDEGP HCRELQGNCSRLGCQHHCVPTLDGPTCYCNS SFQLQADGKTCKDFDECSVYGTCSQLCTNTD GSFICGCVEGYLLQPDNRSCKAKNEPVDRPP VLLIANSQNILATYLSGAQVSTITPTSTRQTTA MDFSYANETVCWVHVGBAAQTQLKCARM PGLKGFVDEHTINISLSLHHVEQMAIDWLTGN FYFVDDIDDRIFVCNRNGDTCVTLLDLELYNP KGIALDPAMGKVFFTDYGQIPKVERCDMDG QNRTKLVDSKIVFPHGITLDLVSRLVYWADA YLDYIEVVDYEGKGRQTIIQGILIEHLYGLTVF ENYLYATNSDNANAQQKTSVIRVNRFNSTEY QVVTRVDKGGALHIYHQRRQPRVRSHACEN DQYGKPGGCSDICLLANSHKARTCRCRSGFS LGSDGKSCKKPEHELFLVYGKGRPGIIRGMD MGAKVPDEHMIPIENLMNPRALDFHAETGFI YFADTTSYLIGRQKIDGTERETILKDGIHNVE GVAVDWMGDNLYWTDDGPKKTISVARLEK AAQTRKTLIEGKMTHPRAIVVDPLNGWMYW TDWEEDPKDSRRGRLERAWMDGSHRDIFVT SKTVLWPNGLSLDIPAGRLYWVDAFYDRIETI LLNGTDRKIVYEGPELNHAFGLCHHGNYLFW TEYRSGSVYRLERGVGGAPPTVTLLRSENPPI

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						WTDWVRRAVQRANKHVGSNMKLLRVDIPQ QPMGIIAVANDTNSCELSPCRINNGGCQDLCL LTHQGHVNCSCRGGRILQDDLTCRAVNSSCR AQDEFECANGECINFSLTCDGVPHCKDKSDE KPSYCNSRRCKKTFRQCSNGRCVSNMLWCN GADDCGDGSDEIPCNKTACGVGEFRCRDGTC IGNSSRCNQFVDCEDASDEMNCSATDCSSYF

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SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence			914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
				amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
	1		1	peptide		/=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion
						RLGVKGVLFQPCERTSLCYAPSWVCDGAND
						CGDYSDERDCPGVKRPRCPLNYFACPSGRCIP
						MSWTCDKEDDCEHGEDETHCNKFCSEAQFE
			i	1		CQNHRCISKQWLCDGSDDCGDGSDEAAHCE
						GKTCGPSSFSCPGTHVCVPERWLCDGDKDCA
						DGADESIAAGCLYNSTCDDREFMCQNRQCIP
						KHFVCDHDRDCADGSDESPECEYPTCGPSEF
						RCANGRCLSSRQWECDGENDCHDQSDEAPK
			l	1		NPHCTSPEHKCNASSQFLCSSGRCVAEALLCN
						GQDDCGDSSDERGCHINECLSRKLSGCSQDC
						EDLKIGFKCRCRPGFRLKDDGRTCADVDECS
			1			TTFPCSQRCINTHGSYKCLCVEGYAPRGGDP
			1			HSCKAVTDEEPFLIFANRYYLRKLNLDGSNY
				}		TLLKQGLNNAVALDFDYREQMIYWTDVTTQ
						GSMIRRMHLNGSNVQVLHRTGLSNPDGLAV
			i			DWVGGNLYWCDKGRDTIEVSKLNGAYRTVL
						VSSGLREPRALVVDVQNGYLYWTDWGDHSL
						IGRIGMDGSSRSVIVDTKITWPNGLTLDYVTE
						RIYWADAREDYIEFASLDGSNRHVVLSQDIPH
J						IFALTLFEDYVYWTDWETKSINRAHKTTGTN
			1			KTLLISTLHRPMDLHVFHALRQPDVPNHPCK
						VNNGGCSNLCLLSPGGGHKCACPTNFYLGSD
						GRTCVSNCTASQFVCKNDKCIPFWWKCDTE
						DDCGDHSDEPPDCPEFKCRPGQFQCSTGICTN
			J			PAFICDGDNDCQDNSDEANCDIHVCLPSQFK
						CTNTNRCIPGIFRCNGQDNCGDGEDERDCPE
						VTCAPNQFQCSITKRCIPRVWVCDRDNDCVD
			1			GSDEPANCTQMTCGVDEFRCKDSGRCIPARW
						KCDGEDDCGDGSDEPKEECDERTCEPYQFRC
						KNNRCVPGRWQCDYDNDCGDNSDEESCTPR
			[			PCSESEFSCANGRCIAGRWKCDGDHDCADGS
						DEKDCTPRCDMDQFQCKSGHCIPLRWRCDA
		-				DADCMDGSDEEACGTGVRTCPLDEFQCNNT
				•		LCKPLAWKCDGEDDCGDNSDENPEECARFV
						CPPNRPFRCKNDRVCLWIGRQCDGTDNCGD
			[			GTDEEDCEPPTAHTTHCKDKKEFLCRNQRCL
			1			SSSLRCNMFDDCGDGSDEEDCSIDPKLTSCAT
						NASICGDEARCVRTEKAAYCACRSGFHTVPG
						QPGCQDINECLRFGTCSQLCNNTKGGHLCSC
						ARNFMKTHNTCKAEGSEYQVLYIADDNEIRS
						LFPGHPHSAYEQAFQGDESVRIDAMDVHVKA
						GRVYWTNWHTGTISYRSLPPAAPPTTSNRHR
						RQIDRGVTHLNISGLKMPRGIAIDWVAGNVY
						WTDSGRDVIEVAQMKGENRKTLISGMIDEPH
						AIVVDPLRGTMYWSDWGNHPKIETAAMDGT
						LRETLVQDNIQWPTGLAVDYHNERLYWADA
]				]		KLSVIGSIRLNGTDPIVAADSKRGLSHPFSIDV
						FEDYIYGVTYINNRVFKIHKFGHSPLVNLTGG
						LSHASDVVLYHQHKQPEVTNPCDRKKCEWL
						CLLSPSGPVCTCPNGKRLDNGTCVPVPSPTPP
						PDAPRPGTCNLQCFNGGSCFLNARRQPKCRC
						QPRYTGDKCELDQCWEHCRNGGTCAASPSG
						MPTCRCPTGFTGPKCTQQVCAGYCANNSTCT
		ļ				VNQGNQPQCRCLPGFLGDRCQYRQCSGYCE
			[			NFGTCQMAADGSRQCRCTAYFEGSRCEVNK
						CSRCLEGACVVNKQSGDVTCNCTDGRVAPS
	1	1	[			CLTCVGHCSNGGSCTMNSKMMPECQCPPHM
1			1			TGPRCEEHVFSQQQPGHIASILIPLLLLLLVL
,						
						VAGVVFWYKRRVQGAKGFQHQRMTNGAM NVEIGNPTYKMYEGGEPDDVGGLLDADFAL

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374	1724	A	3187	191	1815	CLELASAGKIPEESKALSLLAPAPTMTSLMPG AGLLPIPTPNPLTTLGVSLSSLGAIPAAALDPNI ATLGEIPQPPLMGNVDPSKIDEIRRTVYVGNL NSQTTTADQLLEFFKQVGEVKFVRMAGDET QPTRFAFVEFADQNSVPRALAFNGVMFGDRP LKINHSNNAIVKPPEMTPQAAAKELEEVMKR VREAQSFISAAIEPGWLHSTSLCNDFLGCF*RR RMYRE*APCTICGTFHLCLIINWDL*LF*AYTA K*FFPPRVWKEQ*KKRR\RSRSHTRSKSRSSK SHSRKRSQSKHRSRSHNRSRSQKDRRRSK SPHKKRSKSRERRKSRSSSFSHSRDKRKDTREKI KEKERVKEKDREKEREKEKEKEKEKEKEK KDRDKEREKDREKDKEKDREREKEHEKD RDKEKEKEQDKEKEREREKERRRSRSSSRS PRTSKTIKRKSSRSPSPRSRNKKDKKREKERD HISERRERERSTSMRKSSNDRDGKEKLEKNST S
375	1725	A	3192	415	101	AHSSHQTRAILQEFQWDIIRHPPL\SPNLALSG F\FPNLKKSLRGTHFSSVKK\TTLTWLNSQDP WF/FFYP*SPDLQIPSSFRNGLNDWYHHSQKC PDLDGAYVKK
376	1726	A	3199	931	418	GV*WCDLGSPQPPPPGFKQFCLGRSSSWDYR HVPPHPANFVFLLETGFLHAGQAGL\GDPPAS ASQSAGITGVSHTWPKNHLIFYACLVIRSKRI K
377	1727	A	3201	274	1285	KTGYTSRGSPLSPQSSIDSELSTSELEDDSISM GYKLQDLTDVQIMARLQEESLRQDYASTSAS VSRHSSSVSLSSGKKGTCSDQEYDQYSLEDEE EFDHLPPPQPRLPRCSPFQRGIPHSQTFSSIREC RRSPSSQYFPSNNYQQQQYYSPQAQTPDQQP NRTNGDK/PPKKYA*PSPDAKYNCH***QH\SSP VTVRNSQSFDSSLHGAGNGISRIQSCIPSPGQL QHRVHSVGHFPVSIRQPLKATAYVSPTVQGSS NMPLSNGLQLYSNTGIPTPNKAAASGIMGRS ALPRPSLAINGSNLPRSKIAQPVRSFLQPPKPL SSLSTLRDGNWRDGCY
378	1728	A	3202	112	1789	VPGVTESRPSVLRGDHLFALLSSETHQEDPIT YKGFVHKV\ELDRVKLSFSMSLLSRFVGWG* PFKVNFY/TFNRQPLRV\QHRALELTGRWLLW PMLFP\VAPRDVPLLPSDVKLKLYDRSLESNP EQLQAMRHIVTGTTRPAPYIIFGPPGTGKTVT LVEAIKQVVKHLPKAHILACAPSNSGADLLC QRLRVHLPSSIYRLLAPSRDIRMVPEDIKPCCN WDAKKGEYVFPAKKKLQEYRVLITTLITAGR LVSAQFPIDHFTHIFIDEAGHCMEPESLVAIAG LMEVKETGDPGGQLVLAGDPRQLGPVLRSPL TQKHGLGYSLLERLLTYNSLYKKGPDGYDPQ FITKLLRNYRSHPTILDIPNQLYYEGELQACA DVVDRERFCRWAG\LPRQGFPIIFHGVMGKD EREGNSPSFFNPEEAATVTSYLKLLLAPSSKK GKARLSPRSVGVISPYRKQVEKIRYCITKLDR ELRGLDDIKDLKVTCCSTVTPCLPCAPTCPLP ETSSSFHSSPRPRPTPAALNRARALPEPLTPGD
379	1729	A	3206	432	130	SNLRVWDGIRKPACLTNTSCHS PKAAPSVXLWFPPFL*GSFKPTKGHTXCVXIK *LSTREAXDSXPGRQIAXXRQGGKVETTTAL

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						PEQRDHPAPHKQIEAGQGLPGPQAWGG*KGP AXLLPGPGGGPGPVASLEARAQASSGVTPNG GGRTYPYPTFSSGE
381	1731	A	3225	1	840	GTRPGHLPAPSDGFCV/HL*SIPSWGSF*GESL/ EMQLITSLGLQEFDIARNVLELIYAQTLVWIGI FFCPLLPFIQMIMLFIMFYSKNISLMMNFQPPS KAWRASQMMTFFIFLLFFPSFTGVLCTLAITI WRLKPSADCGPFRGLPLFIHSIYSWIDTLSTRP GYLWVVWIYRNLIGSVHFFFILTLIVLIITYLY WQITEGRKIMIRLLHEQIINEGKDKMFLIEKLI KLQDMEKKANPSSLVLERREVEQQGFLHLGE HDGSLDLRSRRSVQEGNPRA
382	1732	A	3238	256	38	LLMIKVSSTCFSCHLHHHHHHHHHHHQGHNS LFFSLKSSSNSSTLPVYLSYNIILVFSKCLVFDF LFSNACL
383	1733	A	3241	1542	343	KGAPSFVRLYQYPNFAGPHAALANKSFFKAD KVTMLWNKKATAVLVIASTDVDKTGASYYG EQTLHYIATNGESAVVQLPKNGPIYDVVWNS SSTEFCAVYGFMPAKATIFNLKCDPVFDFGTG PRNAAYYSPHGHILVLAGFGNLILQI*AD/IMK VWNVKNYKLISKPVASDSTYFAWCPDGEHIL TATCAPRLRVNNGYKIWHYTGSILHKYDVPS NAELWQVSWQPFLDGIFPAKTITYQAVPSEVP NEEPKVATAYRPPALRNKPITNSKLHEEEPPQ NMKPQSGNDKPLSKTALKNQRKHEAKKAAK QEARSDKSPDLAPTPAPQSTPRNTVSQSISGDP EIDKKIKNLKKKLKAIEQLKEQAATGKQLEK NQLEKIQKETALLQELEDLELGI
384	1734	A	3242	3	678	IRSPAARSPGLETPTCLLFVIAAIAAVFVDSAIP RLTQHRPQDGSFPYTILDPPLYLPGQCAPPQP LSQCARRVHGEKLRRPTFGPRHRGAGTAKMS ASLVRATVRAVSKRKLQPTRAALTLTPSAVN KIKQLLKDKPEHVGVKVGVRTRGCNGLSYTL EYTKTKGDSDEEVIQDGVRVFIEKKAQLTLL GTEMDYVEDKLSSEFVFNNPNIKGTCGCGES FNI
385	1735		3243	3190	664	VAMGTPRAQHPPPPQLLFLILLSCPWIQGLPL KEEEILPEPGSETPTVASEALAELLHGALLRR GPEMGYLPGPPLGPEGGEEETTTTIITTTTTT TVTSPVLCNNNISEGEGYVESPDLGSPVSRTL GLLDCTYSIHVYPGYGIEIQVQTLNLSQEEELL VLAGGGSPGLAPRLLANSSMLGEGQVLRSPT NRLLLHFQSPRVPRGGGFRIHYQAYLLSCGFP PRPAHGDVSVTDLHPGGTATFHCDSGYQLQG EETLICLNGTRPSWNGETPSCMASCGGTIHNA TLGRIVSPEPGGAVGPNLTCRWVIEAAEGRRL HLHFERVSLDEDNDRLMVRSGGSPLSPVIYDS DMDDVPERGLISDAQSLYVELLSETPANPLLL SLRFEAFEEDRCFAPFLAHGNVTTTDPEYRPG ALATFSCLPGYALEPPGPPNAIECVDPTEPHW NDTEPACKAMCGGELSEPAGVVLSPDWPQS YSPGQDCVWGVHVQEEKRILLQVEILNVREG DMLTLFDGDGPSARVLAQLRGPQPRRRLLSS GPDLTLQFQAPPGPPNPGLGQGFVLHFKEVPR NDTCPELPPPEWGWRTASHGDLIRGTVLTYQ CEPGYELLGSDILTCQWDLSWSAAPPACQKI

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386	1736	A	3250	5725	3984	GTSTVTMATKKHFSIILNLLGMLLKKDNQDT RKLLMTWALEVAVVMKKSETYAPLFCLPSF HKFCKGLLADTLVEDVNICLQACSSLHALSSS LPDDLLQRCVDVCRVQLVHRGTCIRQAFGKL LKSIPLGVFLSNNNHTEIQEISLALRSHMSKAP SNTFHPQDFSD/VISFILYGNSHRTGKDNWLE RLFYSCQRLDKRDQSTIPRNLLKTDAVLWQW AIWEAAQFTVLSKLRTPLGRAQDTFQTIEGIIR SLAGHTLNPDQDVSQWTTADNDEGHGNNQL RLVLLLQYLENLEKLMYNAYEGCANALTSPP KVIRTFLYTNRQTCQDWLTRIRLSIMRVGLLA GQPAVTVRHGFDLLTEMKTTSLSGNELEVSI MMVVEALCELHCPEAIQGIAVWSSSIVGKHL LWINSVAQQAEGRFEKASVEYQEHLCAMTG VDCCISSFDKSVLTLASAGCKSASLKHCLNGE SRKSVLSKPTDSSPEVINYLGNKACECYISTA DWAAVQEWQNAIHDLKKSTSSTSLNLKADF NYIKSLSSFESGKFVECTEQLELLPGENINLLA GGSKEKIDMKKLLRNM
387	1737	A	3255	380	76	MDIFLYNCKYQVQTEI*NSIQHIMA\SKKLSRF LKYVHNL*AENYKTLMK*INEDLNKQRDVPY S*TARLNKMSIPTKTIFRFKAIYIKIPATYFIET NMQ
388	1738	A	3260	685	428	PQWLGLQVYALPPANFVFFVEMRSTILAQTG FELLDSSDLPASASKSAGITCMSHHARTLSLK *WPFCLSATQEKFC*PASEGVAW
389	1739	A	3269	1	332	LDGYHTPIYMLNRIIRLPAAL*IISDQTGHALTI LTRLETQMINADYQNKLTLDYLLTTDREVYE PFNLTNYCLHIHNQRLGAYDLG*V*Q/KLAHV PVQV*HGFDPEAMFR
390	1740	A	3270	2	372	GRCHDQNKGKS\DGPDAQAEACGGESTYQEL LVNQNPIGQPLACRRLTRKIYEGIKKAVKPNH SPRGVKKVHKFVNKGEKGIMVLAGDTLGIGV YCLLPCMC*DRKLTYAHIPSTTDLGAGAGY
391	1741	A	3273	1	187	FFQEMLDIMKAISDMMGKCTYPVLKEDAPRQ HVETFFQIEELTRSQEGMKLGENFLMFAMPP DDSKESKGK*FFQEMLDIMKAISDMMGKCTY PVLKEDAPRQHVETFFQVGINQKSRGHEVRR KFPDVCHAPR
392	1742	A	3281	901	521	FFFGDGVSPCRQAGV*WHDLDSLQNLPPGFK RFSYLSI.PSSW\DYRHVLPRQANFCIF/M*RRG FTMLARMVSIS*PRDLPALASQSAGITGVSHH APPQMDFTFALLCFALKGCLPRQKEGGTLNLI
393	1743	A	3283	385	3	RNRSVVPEFVLLGLSAGPQTQTLLFVLFVVIC LLTVMGNLLLLVVINADSCLHTPMYFFLGQL SFLDLCHSSVTAPKLLENLLSEKKTISVEGCM A*VFFVFATGGTESSLLAVMAYDRYVAIRTR G
394	1744	Ā	3284	575	1054	CTKCKADCDTCFNKNFCTKCKSGFYLHLGKC LDNCPEGLEANNHTMECVSIVHCEVSEWNP WSPCTKKGKTCGFKRGTETRVREIIQHPSAKG NLCPPTNETRKCTVQRKKCQKGERGKKGRE

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395	1745	A	3286		340	KQQQ RVLYVPSMGFCILVAHGWQKISTKSVFKKLS
393		A		1		WICLSMVILTHSLKTFHRNWDWESEYTLFMS ALKVNKNNAKLWNNVGHALENEKNFERAL KYFLQATHVQPDDIGAHMNVGR
396	1746	A	3293	1	172	GFRAVVMTVKTEAAKGTLTYSRMRGMVAIL IAFMKQRRMGLNDFIQKIANNSYACKO
397	1747	A	3295	12	401	AEPACGASSCTPPSLRSSSSQSVGPLRPGRPL WSEACAFL*AAAPQGPASPCCGLPSGFPRVW AQCCPPGGALRFPEGLGSVLSPRRCPQVSRGS GLSAVPQEVPSGFLGPGLRACPQEAPSRFLRA GLT
398	1748	A	3300	1912	2768	KQRRWQNIQRKGPKRYIVIAGNSQSHQPMIFS MLRKLPKVTCRDVLPEIRAICIEEIGCWMQSY STSFLTDSYLKYIGWTLHDKHREVRVKCVKA LKGLYGNRDLTARLELFTGRFKDWMVSMIV DREYSVAVEAVRLLILILKNMEGVLMDVDCE SVYPIV*ASN*GLASAVGEFLYWKLFYPECEI RTMGGREQRQSPGAQRTFFQLLLSFFVESKSH SVTQAGVQWQFSAHRDLCLPGSSNSHVSASR VAGIAGAHRHTWLIYVFFSWRQGFAVLAGL VSNS
399	1749	A	3301	536	2391	LRSYGCKAPSRISHLHK\FLFLLLPSLLMGYSE SPPPITDSWAPFISLTHHVLSQSQSPLSSNCWI CLSTHTQ*FTALPADLLTWTQSNVSLHISYLAI PFLADSFLKPV/L*PGNSAKHLSFKLSSLSMVS GRAVALLHLIASGLTSIQTNTASSKPPIWGYL STQTSFISPPPLCLSRTYPNPAHATMVGQVPQ SLCGLIFTL/RTPCRPSILHPNYKIISTSAWQKV LCFSGSPTIHTSLHLTTGSSFLSFHPIPGFPAAN SALYVSSLKGPPGKNVTIPSPVTGT*QPPHRGS N/RLTVDKDNFFLSPKPNSLHQLPSQ\TPYQAL TGAALAGSYPIWENENTLSWLPTFTYNFCLST PSLFFLCDTN*YLCLPANWSGTCTLVFQAPTI NILPPNQTILISVEASISSSPIRNKWALHLITLLT GLGITAALGTGIAGITTSITSYQTLFTTLSNTVE DMHTSITSLQRQLDFLVGVILQNWRVLDLLT TEKGGTCIYLQEECCFCVNESGIVHIAVRRLH DRAAEL*HQVADSWWQGSSLLRWIPWVAPF LGPLIFLFLLLMIGPCIFNLVSRFISQRLNCFIQ ASMQKHIDNIFHLCHV*YQSLRGNHSEAPEPR P
400	1750	A	3303	2	453	THWRHSSGVPGSTTARRRRRELEIATSDNQE YYNRLCQEVTNRERNDQKMLADLDDLNRTK KYLEERLIELLRDKDALWQKSDALEFQQKLS AEERWLGDTEANHCLDCKREFSWMVRRHHC RICGRIFCYYCCNNYVLSKHGGKKERCC
401	1751	A	3304	1	626	MAPQHSSLDDKVPQQASTVCFEFQDILQHSQ CTEHKDSLWGPGARSQPFGAHNTRLSPDSCP EKIVLRALKDSRAGMPEQDKDPGVQENPDD QRRVPQGTGDAPSAFRPLWDNGGLSPFVSRP GPLERDLHAQRSEVTYNQRSQSSWMSSFPKR NAFVSPYSSMGQAQP/GLPKTNPIGESCCWEG LSLSTQILG*QKPSKYIPSLCKR
402	1752	A	3305	1678	172	MELPSGPGPERLFDSHRLPGDCFLLLVLLLYA PVGFCLLVLRLFLGIHVFLVSCALPDSVLRRF VVRTMCAVLGLVARQEDSGLRDHSVRVLISN HVTPFDHNIVNLLTTCSTVSESEAESATGRFP

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  GAQLKAPLSPLAFRMEDTEALPLTPILYPTCQ FFFF\IFLLAFSSPGSQPLLNSPPSFVCWSR GFMEMNGRGELVESLKRFCASTRLPPTPLLLF PEEEATNGREGLLRFSSWPFSIQDVVQPLTLQ VQRTLVSVTVSDASWVSELL\WSLFVPFTVY QVRWLRPVHRQLGEANEEFALRVQQ\LVAKE
						LG\QTGTRLTPA\DKAEHMKRQRHPR\LRPQS AQSSFPPSPWVLSS/SDVQTGQTLGFREFKESF CPHVAIGVFIPERPWPKTGCCKTLTIHLILL*G GPVSFSCPE\DIHPRGT*VPTQQASGLPSFPSYG PARGGVL*HPSAQQPLTFA\KSS\WARAGRAL QERKQ\ALYEYARRRFTERRAPGGLD
403	1753	A	3307	44	447	DPSPSLLAVALGLRAGERTRSGPGSSSPSGGIS GGASAGLASSPECACGRSHFTCAVSALGECT CIPAQWQCDGDNDCGDHSDEDGCILPTCSPL DFHCDNGKCIRRSWVCDSDNDCEDDSDEQD CPPRECEED
404	1754	A	3311	409	1	PRHGWGRRVLGRDRPRLQKVKKSVKAIYIPG QDIIVQNEEIYARVLDKFGSNFLSRDNADLGT AFVKFSTLTK*LSALLKNLLQGLSRNVIFTLDS LLKGDLKGVKGDLKKPFDKAWKDYETKFAK IEKEKREREWR
405	1755	A	3322	12	458	AAVPVENPWDDPRVRPRVRIFTWEDCIAGQA KVLCNDSYGVTIDWSPKGAFIRLTSQSVGNG HPASKENDQMVDTIKNTTKVPIIWTYGDMVE PRPQMIRPAVGAKHKELWKILMALKKIK\IWE GKYTKPSQYNPNYMLELAHNDSVW
406	1756	A	3324	1	426	LSMLSTISTEHRLSVLWPIWYCCHCPTHLSAV MCVLLWALSLLQSILEWMFCSFLFSDVDSDN WCQILDFLTAVWLIFLILVLCGFTLVLLVRIIC GSQKMPLTRLYVTILLTGLVFLFCSLPLSIQ*F LLYWIEKDLDDL
407	1757	A	3328	213	1841	SGDLSPAELMMLTIGDVIKQLIEAHEQGKDID LNKVKTKTAAKYGLSAQPRLVDIIAAVPPQY RKVLMPKLKAKPIRTASGIAVVAVMCKPHRC PHISFTGNICVYCPGGPDSDFEYSTQSYTGYEP TSMRAIRARYDPFLQTRHRIEQLKQLGHSVD KVEFIVMGGTFMALPEEYRDYFIRNLHDALS GHTSNNIYEAVKYSERSLTKCIGITIETRPDYC MKRHLSDMLTYGCTRLEIGVQSVYEDVARD TNRGHTVKAVCESFHLAKDSGFKVVAHMMP DLPNVGLERDIEQFTEFFENPAFRPDGLKLYP TLVIRGTGLYELWKSGRYKSYSPSDLVELVA RILALVPPWTRVYRVQRDIPMPLVSSGVEHG NLRELALARMKDLGIQCRDVRTREVGIQEIH HKVRPYQVELVRRDYVANGGWETFLSYEDP DQDILIGLLRLRKCSEETFRFELGGGVSIVREL HVYGSVVPVSSRDPTKFQHQGFGMLLMEEA ERIAREEHGSGKIAVISGVGTRNYYRKIGYRL QGPYMVKMLK AIASPRAAGIRHELTSTMAAGKNKRLTKGGK
400	1/38	A	3333	3	40/	AIASPRAAGIRHELTSTMAAGKNKRETKGGK KGAKKKAV/DNIINIGKTLVTRTQRTKIASDG LKGRVFEESLADLQND\TDGYLLRVI*VAFTT ERTNQI/REVFNKLIPDSIGKDIEKACQSIYPLH DDFARKVKMLKKPKFELRKLMELHGEGSS
409	1759	Α	3338	7	1252	PRWRNSARDEILLSFPQNYYIQWLNGSLIHGL WNLASLFSNLCLFVLMPFAFFFLESEGFAGLK KGIRARILETLGMLLLLALLILGIVWVASALID NDAASMESLYDLWEFYLPYLYSCISLMGCLL

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410	1760	A	3339	127	1433	AL GSHRFSLASPLDPEVGPYCDTPTMRTLFNLL WLALACSPVHTTLSKSDAKKAASKTLLEKSQ FSDKPVQDRGLVVTDLKAESVVLEHRSYCSA KARDRHFAGDVLGYVTPWNSHGYDVTKVFG SKFTQISPVWLQLKRRGREMFEVTGLHDVDQ GWMRAVRKHAKGL\P*CLGSCLRTGLTMISG/ YVLDSEDEIEELSKTVVQVAKNQHFDGFVVE VWNQLLSQKRVGLIHMLTHLAEALHQARLL ALLVIPPAITPGTDQLGMFTHKEFEQLAPVLD GFSLMTYDYSTAHQPGPNAPLSWVRACVQV LDPKSKWRSKILLGLNFYGMDYATSKDAREP VVGARYIQTLKDHRPRMVWDSQVSEHFFEY KKSRSGRHVVFYPTLKSLQVRLELARELGVG VSIWELGQGLDYFYDLL*VGIAASAVDVFFSK PWSE
411	1761	A	3342	74	2701	VATRKLAKGFTQFAKMTEGTKKTSKKFKFK FKGFGSFSNLPRSFTLRRSSASISRQSHLEPDTF EATQDDMVTVPKSPPAYARSSDMYSHMGTM PRPSIKKAQNSQAARQAQEAGPKPNLVPGGV PDPPGLEAAKEVMVKATGPLEDTPAMEPNPS AVEVDPIRKPEVPTGDVEERPPRDVHSERAA GEPEAGSDYVKFSKEKYILDSSPEKLHKELEE ELKLSSTDLRSHAWYHGRIPREVSETLVQRN GDFLIRDSLTSLGDYVLTCRWRNQALHFKIN KVVVKAGESYTHIQYLFEQESFDHVPALVRY HVGSRKAVSEQSGAIIYCPVNRTFPLRYLEAS YGLGQGSSKPASPVSPSGPKGSHMKRRSVTM TDGLTADKVTRSDGCPTSTSLPRPRDSIRSCA LSMDQIPDLHSPMSPISESPSSPAYSTVTRVHA APAAPSATALPASPVARRSSEPQLCPGSAPKT HGESDKGPHTSPSHTLGKASPSPSLSSYSDPDS GHYCQLQPPVRGSREWAATETSSQQARSYGE RLKELSENGAPEGDWGKTFTVPIVEVTSSFNP ATFQSLLIPRDNRPLEVGLLRKVKELLAEVDA RTLARHVTKVDCLVARILGVTKEMQTLMGV RWGMELLTLPHG\RKLRLDLLERFHTMSIML AVDILGCTGSAEERAALLHKTIQLAAELRGT MGNMFSFAAVMGALDMAQISRLEQTWVTLR QRHTEGAILYEKKLKPFLKSLNEGKEGPPLSN TTFPHVLPLITLLECDSAPPEGPEPWGSTEHGV EVVLAHLEAARTVAHHGGLYHTNAEVKLQG FQARPELLEVFSTEFQMRLLWGSQGASSSQA RRYEKFDKVLTALSHKLEPAVRSSEL
412	1762	A	3347	1	898	RRYEKFDKVLTALSHKLEPAVRSSEL  IDRAAECRTKPLPMAVSIRGNADSIVACLVLM VLYLIKKRLVACAAVFYGFAVHMKIYPETYI LPITLHLLPDRDNDKSLRQFRYTFQACL*ELL KRLCNRTALMFVAVAGLTFFALSFGFYYEYG WEFLEHTYFYHLTRRDIRHNFSPYFYMLYLT AESKWSFSLGIAAFLPQLILLSAVSFAYYRDL VFCWFLHTSIFVTFNKVCTSQYFLWYLCLLPL

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413	1763	A	3361	3	474	SHYKEEPLTERIKYD PIPVRWNSLEGRLLRGYEQHANDGKDYISRN *DLRSWTAADMAAQITKRKWEAEEFAEQIKA YLEGTCVER/LRTHLENGKETLQLTEQSSQPTI PIVGIVAGLVLLGAVVTGAVVSAVMCRKKNS GHFLPTDRVSYSEAASSDHAQGSDVSLTACK V
414	1764	A	3363	1488	453	HQILELKKKILKTYNPDYDEDLVQEASSEDVL GVHMVDKDTERDIEMKRQLRRLRELHLYST WKKYQEAMKTSLGVPQRERDEGSLGKPLCP PEILSETLPGSVKKRVCFPSEDHLEEFIAEHLP EASNQSLLTVAHADAGTQTNGDLEDLEEHGP GQTVSEBATEVHMMEGDPDTLAELLIRDVLQ ELSSYNGEEE\DPEEVKTSLGVPQRGDLEDLE EHVPGQTVSEBATGVHMMQVDPATLAKSDL EDLEEHVPEQTVSEBATGVHMMQVDPATLA KQLEDSTITGSHQQMSASPSSAPAEEATEKTK VEEEVKTRKPKKKTRKPSKKSRWNVLKCWD IFNIF
415	1765	A	3369	431	315	IPWSWVGRLSVRKMSILF*LTYNYNAILNKTP PSFSPSL
416	1766	A	3373	42	651	RQEKMGLGEIGASGVLRSMLKERKKQNMKG NGNVTLTPLLPAVQCGCHLQPAGRSPLPSSHS APGLCSPLHPLQPQQEASTCPSGTLQGREKAA PGQGRPLCSLWAGGAGA\PGERGAEGRGPSD QAPDPKSGPWLFPPGLGAPAEVRLHNVPHNL RRPPLP*ARGK*PPNSGCPWSEGRAKQPLSCG PKPQCSLPSQVPGDTH
417	1767	A	3382	2	2061	EAQDPRACGPDAGGRFAARDAPGNSLRPPPS SPP/GWPGQLRLLPRVPGSELRCGKPERGRLP ASPPGKIRGWPPGISKRPGLGGRSFPPGFAPRT WRPEARGPSVQSLPPIFSPQSAQTTAR*RPGAP KNAGRCGGA\RGPRLSLGPPPGPPPAPALPAR ASAGAGAAAALAVGGVRGAGGARGTGGY GHCSGR/PTGRTGPGPQGPGPPMPARPR*AS\S TRGSRRGPGSRPARAAAAPRAGDHGRRPVRV HLRQHTAV*EPRLGDATAPPGGAAGPGAPAP R\GPGWDCALLPSPGPRSPRAVGCAEPEIWDP SPRRGTSPVPSVRSLRSEPANPRLGLPALLNSY PLKGPGLPPPWGPRTQTGHVIITVQPSGSCIEH SKSLD/RGPWGAPPWGPSSSGLCSPKLATAGP PQSWGLCQIGRRRGLGGPGLKRGET/GLL*GC SMDHANRTKGPGVPTSNRCFSHIPG\GDGCSD HSSCEGHPDLHAGREMPAAPGLSELERVRFT VGCGGLASGISSASVSGLSPNRAGGPGQGDW EMYPVSWQTQESGGQG/SPKTGR*VGMLQA GAGSLQGTGDGVWGLWEDGP/RG*DSPLPS GTGTEP*TPTTSIPFFPQPSGVYPSRATLLPMPS Y*ALGPSANKSEKPLLSFLYRGLCCRISLQLA KGIGQLSEIPLLNVETAFWSMWVTYFRK
418	1768	A	3398	304	2121	EEEEEEDEDDDDNNEEEEFECYPPGMKVQV RYGRGKNQKMYEASIKDSDVEGGEVLYLVH YCGWNVRYDEWIKADKIVRPADKNVPKIKH RKKIKNKLDKEKDKDEKYSPKNCKPPALGPN PPFQTNPISWKWYPKLDLTDAKNSDTAHIKSI EITSILNGLQASESSAEDSEQEDERGAQDMDN NGKEESKIDHLTNNRNDLISKEEQNSSSLLEE

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first	Predicted end nucleotide location corresponding to last amino acid residue	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
denes				amino acid residue of peptide sequence	of peptide sequence	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
						NKVHADLVISKPVSKSPERLRKDIEVLSEDTD YEEDEVTKKRKDVKKDTTDKSSKPQIKRGKR RYCNTEECLKTGSPGKKEEKAKNKESLCMEN SSNSSSDEDEETKAKMTPTKKYNGLEEKRK SLRTTGFYSGFSEVAEKRIKLLNNSDERLQNS RAKDRKDVWSSIQGQWPKKTLKELFSDSDTE AAASPPHPAPEEGVAEESLQTVAEESCSPSV ELEKPPPVNVDSKPIEEKTVEVNDRKAEFPSS GSNFSA*IPLPYLHLNRLHQSL*QKGSRQQSS VTVSEPLAPNQEEVRSIKSETDSTIEVDSVAGE LQDLQSERE*LASRF*CQCELKQ**SARTRTS* KSLYRSEKSERCSGRRKFIKKAEKKP*SNSGK QQKEGK
419	1769	A	3399	206	463	QRECLSIHIGQAGIQIGDACWELYCLEHGIQP NGVVLDTQQDQLENAKMEHTNASFDTFFCE TRAGKHVPRALFVDLEPTVIDGIR
420	1770	A	3408	1010	685	RRLSFFF*IWSSVLVTQARVQWRDLGSPQPLP PGFKRFSCLSLPSSWDYRHPSPRPVNF/HVFLV VMGFHHVGQAGLELLTSGDLPALASQSARIT GVNHCAQPRGHFH
421	1771	A	3409	355	1326	ADSNLIESCWQELGLGPWGGDWRVEQVGAS ASLRFPREVCSIRFLFTAVSLLSLFLSAFWLGL LYLVSPLENEPKEMLTLSEYHERVRSQGQQL QQLQAELDKLHKEVSTVRAANSERVAKLVF QRLNEDFVRKPDYALSSVGASIDLQKTSHDY ADRNTAYFWNRFSFWNYARPPTVILEPHVFP GNCWAFEGDQGQVVIQLPGRVQLSDITLQHP PPSVEHTGGANSAPRDFAVFFLLSFFTHQGLQ VYDETEVSLGKFTFDVEKSEIQTFHLQNDPPA AFPKVKIQILSNWGHPRFTCLYRVRAHGVRT SEGAEGSAQGPH
422	1772	A	3412	2	421	EFDAQPSIGALVVFKRP*ATTGSDPGPKRGMN YLVSCSMRSPESGKGEPGTARDYTPMGRPPP PVPSVSPGPLPGSLAIAPHSPEPHPWEQQPPRG QARSPPGGWLGSAT/RVRRPHNHP/RGH/HSP VDTAGAPASPGPDVCE
423	1773	A	3420	91	706	DAQRAIYSSVGPAVSLRQRQQDGAVKESGR/ RGGVRSFSRAAAAMAPIKVGDAIPAVEVFEG EPGNKVNLAELFKGKKGVLFGVPGAFTPGCS KTHLPGFVEQAEALKAKGVQVVACLSVNDA FVTGEWGRAHKAEGKVRLLADPTGAFGKET DLLLDDSLVSIFGNRRLKRFSMVVQDGIVKA LNVEPDGTGLTCSLAPNIISQL
424	1774		3421	4	7688	RQVTRVGTRVLGSTTAAVFLSVEDDNDNAPQ FSEKRYVVQVREDVTPGAPVLRVTASDRDKG SNAVVHYSIMSGNARGQFYLDAQTGALDVV SPLDYETTKEYTLRVRAQDGGRPPLSNVSGL VTVQVLDINDNAPIFVSTPFQATVLESVPLGY LVLHVQAIDADAGDNARLEYRLAGVGHDFP FTINNGTGWISVAAELDREEVDFYSFGVEAR DHGTPALTASASVSVTALDVNDNNPTFTQPE YTVRLNEDAAVGTSVVTVSAVDRDAHSVITY QITSGNTRNRFSITSQSGGGLVSLALPLDYKLE RQYVLAVTASDGTRQDTAQIVVNVTDANTH RPVFQSSHYTVNVNEDRPAGTTVVLISATDE DTGENARITYFMEDSIPQFRIDADTGAVTTQA ELDYEDQVSYTLAITARDNGIPQKSDTTYLEI LVNDVNDNAPQFLRDSYQGSVYEDVPPFTSV LQISATDRDSGLNGRVFYTFQGGDDGDGDFI

NO: of nucl- nucl- nucl- eotide seq- uence  NO: of nucl- peptide eotide seq- uence  NO: of nucl- peptide eotide seq- uence  NO: of nucl- peptide eotide seq- uence  NO: of peptide eotide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  NO: of peptide seq- uence  No: of peptide sequence  No: of peptide s	cine, H=Histidine,  =Leucine, agine, P=Proline, e, S=Serine, W=Tryptophan, n, *=Stop codon, stion, \=possible ENVAQYVLRAYAVDK LDVNDNPPVFEQDEFD TATDPDEGTNAQIMY
sequence  uence  09/496 914  orrespondi ng to first amino acid residue of peptide sequence  versidue of peptide sequence  orrespondi ng to first amino acid residue of peptide sequence  versidue of peptide sequence  orrespondi ng to last amino acid residue of peptide sequence  M=Methionine, N=Aspara Q=Glutamine, N=Aspara T=Threonine, V=Valine, Y=Tyrosine, X=Unknowr /=possible nucleotide delegation in the last amino acid residue of peptide sequence	agine, P=Proline, e, S=Serine, W=Tryptophan, n, *=Stop codon, stion, \=possible ENVAQYVLRAYAVDK LDVNDNPPVFEQDEFD TATDPDEGTNAQIMY
uence  914  ng to first amino acid residue of peptide residue of peptide sequence  914  ng to first amino acid residue of peptide residue of peptide sequence  914  ng to first amino acid residue of peptide residue of peptide sequence  92  Glutamine, R=Arginin T=Threonine, V=Valine, Y=Tyrosine, X=Unknowr /=possible nucleotide dele nucleotide insertion	e, S=Serine, W=Tryptophan, n, *=Stop codon, stion, \=possible ENVAQYVLRAYAVDK LDVNDNPPVFEQDEFD TATDPDEGTNAQIMY
amino acid residue of peptide residue of peptide sequence y=Tyrosine, X=Unknowr /=possible nucleotide dele nucleotide insertion	W=Tryptophan, n, *=Stop codon, stion, \=possible ENVAQYVLRAYAVDK LDVNDNPPVFEQDEFD TATDPDEGTNAQIMY
residue of peptide sequence Y=Tyrosine, X=Unknowr /=possible nucleotide dele nucleotide insertion	n, *=Stop codon, stion, \=possible ENVAQYVLRAYAVDK LDVNDNPPVFEQDEFD TATDPDEGTNAQIMY
peptide /=possible nucleotide dele sequence nucleotide insertion	tion, \=possible ENVAQYVLRAYAVDK LDVNDNPPVFEQDEFD TATDPDEGTNAQIMY
sequence nucleotide insertion	ENVAQYVLRAYAVDK LDVNDNPPVFEQDEFD TATDPDEGTNAQIMY
	LDVNDNPPVFEQDEFD FATDPDEGTNAQIMY
I I I I VESTSGIVKTEKKLUKI	LDVNDNPPVFEQDEFD FATDPDEGTNAQIMY
GMPPARTPMEVTVTVI	
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TGDYCETEVDLCYSRP	
CLCRDGYTGEHCEVSA	
CVNLLVGGFKCDCPSC	
AHSFITFRGLRQRFHFT YNGRFNEKHDFVALEY	
TTVSPFVPGGVSDGQW	HTVOLKYYNKPLLG
QTGLPQGPSEQKVAVV	TVDGCDTGVALRFGS
VLGNYSCAA\QGTQGC	
VPDLPESFPVRMRQFV	GCMRNLQVDSRHIDM
ADFIANNGTVPGCPAK	
VNQWDAFSCECPLGFC	
LGSSLVAWHGLSLPISO	
GVLLQAITRGRSTITLQ QASSLRLEPGRANDGD	
GHAILSFDYGQQRAEG	
GGIPGPAGGVARGFRG	
SLDPSHGESINVEQGCS	
CSNDWDSYSCSCDPGY	YYGDNCTNVCDLNPC
EHQSVCTRKPSAPHGY	,
RIDQPCPRGWWGHPTC	
NKTSGECHCKENHYRE SLSRVCDPEDGQCPCK	
AEVITNGCEVNYDSCF	
LPAAAPCPKGSFGTAV	
NCTSITFSELKGFAERL	
ALLLRNATQHTAGYFO	SDVKVAYQLATRLL
AHESTQRGFGLSATQD	VHFTENLLRVGSALL
DTANKRHWELIQQTEO	
ALAQNMRHTYLSPFTI	
FAGAKLPRYEALRGEQ ETPPVVRPAGPGEAQE	
GEAVASVIIYRTLAGLL	, , ,
RPIINTPVVSISVHDDEE	
LLETEERTKPICVFWNF	`
EVVFRNESHVSCQCNH	
NGEILPLKTLTYVALGY	
RILRSNQHGIRRNLTAA	` `
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PVSGLQPSFAVLLLLSA	
FHYLFATCNCIQGPFIFI	
LACSRKPSPDPALTTKS	
RLYQP\YGDSAGSLHST	SRSGKSQPSYIPFLLR
EESALNPG\QGPPGLGG	
H\DS*TRDFDSDLSLEDI	DQSGSYASTHSSDSEE

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425	1775	A	3429	155		EERLRENGDALSREGSLGPLPGSSAQPHKGIL KKKCLPTISEKSSLLRLPLEQCTGSSRGSSASE GSRGGPPSRPPPRQSLQEQLNGVMPIAMSIKA GTVDEDSSGSEFLFFNFLH
		A			1417	GEPAVQSCDCGCTQRSCPWLLVAPGLLSSSS RAASVREAEDAPLQPASIHPVSQGSRGPEGSL GSAECLPGDPLGARRATRAHSPVPGPPPSLPA AGTAVKRGLQPG*GA/GATSTPGTGAATGGL CGPAWAAPSAVGPCCCCPSISTTPSQMRSARP SLGCLPSWAS\PGTEHPPGPQGPGPS*DLCSV* KREFQRGPWAGMVILHRISAADPARAPGPDS NLQSALQQPATGCSEPAAVYSPPIGLWGA**P EYG*PQHSLPG*TAPADR*P\AGIKDRVYSNSI YELLENGQRAGTCVLEYATPLQTLFAMSQYS QAGFSREDRLEQAKLFCRTLEDILADAPESQN NCRLIAYQEPADDSSFSLSQEVLRHLRQEEKE EVTVGSLKTSAVPSTSTMSQEPELLISGMEKP LPLRTDFS
426	1776	A	3431	1662	369	AIWWLSWLQHDLLPTPTQVAIDFTASNGDPR SSQSLHCLSPRQPNHYLQALRAVGGICQDYD/ SVGESGAGGNRQGGLAQRIPQLFLLPSDKRFP AFGFGARIPPNFEVG*MRGKEGDGGRVSQAE KAGPHCSRLALTG\SHDFAINFDPENPECEGK RGDFHLPRLPADTLHTGAQTPLPRAQLPVPST HPRPVFI\EISGVIASYRRCLPQIQLYGPTNVAP IINRVAEPAQREQSTGQATKYSVLLVLTDGV VSDMAETRTAIVRASRLPMSIIIVGVGNADFS DMRLLDGDDGPLRCPRGVPAARDIVQFVPFR DFKDVSPPGPFRLKDSSASHPPKSDLRLPPFD VLLRTREPSWPP*SPTSPSDDPASPTLPLTPNHI TVPTL\AAPSALAKCVLAEVPRQVVEYYASQ GISPGAPRPCTLATTPSPSP
427	1777	A	3446 .	79	9748	GCQSCWPAWPRLRRRGPASAGARLGRKAPW GLPGRVQDGRPLRFCFYLRPRAPFIAPVLSGA ASRPEASGDCRAGRETAMATLEKLMKAFESL KSFQQQQQQQQQQQQQQQQQQQQQPPPP PPPPPPPPPQLPQPPPPQAQPLLPQPQPPPPPPPPPP

SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-	l	USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence	uciico		914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
uciico			714	amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
				peptide	sequence	/=possible nucleotide deletion, \=possible
						l •
				sequence		nucleotide insertion
			1			TTEYPEEQYVSDILNYIDHGDPQVRGATAILC
		i	i		}	GTLICSILSRSRFHVGDWMGTIRTLTGNTFSL
,						ADCIPLLRKTLKDESSVTCKLACTAVRNCVM
						SLCSSSYSELGLQLIIDVLTLRNSSYWLVRTEL
		1				LETLAEIDFRLVSFLEAKAENLHRGAHHYTGI
						LKLQERVLNNVVIHLLGDEDPRVRHVAAASL
		ļ				IRLVPKLFYKCDQGQADPVVAVARDQSSVYL
						KLLMHETQPPSHFSVSTITRIYRGYNLLPSITD
			[			VTMENNLSRVIAAVSHELITSTTRALTFGCCE
						ALCLLSTAFPVCIWSLGWHCGVPPLSASDESR
		İ	1			KSCTVGMATMILTLLSSAWFPLDLSAHQDAL
						ILAGNLLAASAPKSLRSSWASEEEANPAATK
		l		}		QEEVWPALGDRALVPMVEQLFSHLLKVINIC
			1			AHVLDDVAPGPAIKAALPSLTNPPSLSPIRRK
		İ				GKEKEPGEQASVPLSPKKGSEASAASRQSDTS
•						GPVTTSKSSSLGSFYHLPSYLKLHDVLKATHA
						NYKVTLDLQNSTEKFGGFLRSALDVLSQILEL
			]			ATLODIGKCVEEILGYLKSCFSREPMMATVC
	I					VQQLLKTLFGTNLASQFDGLSSNPSKSQGRA
						QRLGSSSVRPGLYHYCFMAPYTHFTQALADA
		1	İ			SLRNMVQAEQENDTSGWFDVLQKVSTQLKT
		j	j			NLTSVTKNRADKNAIHNHIRLFEPLVIKALKO
			}			YTTTTCVQLQKQVLDLLAQLVQLRVNYCLL
						DSDQVFIGFVLKQFEYIEVGQFRESEAIIPNIFF
						FLVLLSYERYHSKQIIGIPKIIQLCDGIMASGR
						KAVTHAIPALQPIVHDLFVLRGTNKADAGKE
	ĺ	ļ				LETQKEVVVSMLLRLIQYHQVLEMFILVLQQ
			ł			CHKENEDKWKRLSRQIADIILPMLAKQQMHI
		1	1			DSHEALGVLNTLFEILAPSSLRPVDMLLRSMF
				'		VTPNTMASVSTVQLWISGILAILRVLISQSTED
			·			IVLSRIQELSFSPYLISCTVINRLRDGDSTSTLE
						EHSEGKQIKNLPEETFSRFLLQLVGILLEDIVT
		[				KQLKVEMSEQQHTFYCQELGTLLMCLIHIFKS
						GMFRRITAAATRLFRSDGCGGSFYTLDSLNLF
						ARSMITTHPALVLLWCQILLLVNHTDYRWW
						AEVQQTPKRHSLSSTKLLSPQMSGEEEDSDLA
						AKLGMCNREIVRRGALILFCDYVCQNLHDSE
		ĺ .			ľ	HLTWLIVNHIQDLISLSHEPPVQDFISAVHRNS
						AASGLFIQAIQSRCENLSTPTMLKKTLQCLEGI
		1	İ			HLSQSGAVLTLYVDRLLCTPFRVLARMVDIL
						ACRRVEMLLAANLQSSMAQLPMEELNRIQEY
						LQSSGLAQRHQRLYSLLDRFRLSTMQDSLSPS
						PPVSSHPLDGDGHVSLETVSPDKDWYVHLVK
						SQCWTRSDSALLEGAELVNRIPAEDMNAFM
						MNSEFNLSLLAPCLSLGMSEISGGQKSALFEA
						AREVTLARVSGTVQQLPAVHHVFQPELPAEP
		ļ				AAYWSKLNDLFGDAALYQSLPTLARALAQY
						LVVVSKLPSHLHLPPEKEKDIVKFVVATLEAL
						SWHLIHEQIPLSLDLQAGLDCCCLALQLPGL
						WSVVSSTEFVTHACSLIYCVHFILEAVAVQPG
						EQLLSPERRTNTPKAISEEEEEVDPNTQNPKYI
						TÄACEMVAEMVESLQSVLALGHKRNSGVPA
						FLTPLLRNIIISLARLPLVNSYTRVPPLVWKLG
						WSPKPGGDFGTAFPEIPVEFLQEKEVFKEFIYR
					4	INTLGWTSRTQFEETWATLLGVLVTQPLVME
						QEESPPEEDTERTQINVLAVQAITSLVLSAMT
						VPVAGNPAVSCLEQQPRNKPLKALDTRFGRK
					•	LSIIRGIVEQEIQAMVSKRENIATHHLYQAWD
						PVPSLSPATTGALISHEKLLLQINPERELGSMS
						YKLGQVSIHSVWLGNSITPLREEEWDEEEEEE
						L INIANUVAIDAV WIADVALLELKEEEWDEEEEEE

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of, peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  ADAPASSPPTSPVNSRKHRAGVDIHSCSQFL LELYSRWILPSSSARRTPAILISEVVRSLLVVS DLFTERNQFELMYVTLTELRRVHPSEDEILAQ YLVPATCKAAAVLGMDKAVAEPVSRLLESTL RSSHLPSRVGALHGILYVLECDLLDDTAKQLI PVISDYLLSNLKGIAHCVNIHSQQHVLVMCAT AFYLIENYPLDVGPEFSASIIQMCGVMLSGSE ESTPSIIYHCALRGLERLLLSEQLSRLDAESLV KLSVDRVNVHSPHRAMAALGLMLTCMYTG KEKVSPGRTSDPNPAAPDSESVIVAMERVSVL FDRIRKGFPCEARVVARILPQFLDDFFPPQDIM NKVIGEFLSNQQPYPQFMATVVYKVFQTLHS TGQSSMVRDWVMLSLSNFTQRAPVAMATWS LSCFFVSASTSPWVAAILPHVISRMGKLEQVD VNLFCLVATDFYRHQIEEELDRRAFQSVLEV
428	1778	A	3449	3	430	VAAPGSPYHRLLTCLRNVHKVTTC NSRPSPSAALVEVLLRSGSTFPHTVSGGWAA WGPWSSCSRDCELGFRVRKRTCTNPEPRNGG LPCVGDAAEYQDCNPQACPVRGAWSCWTS WSPCSASCGGGHYQRTRSCTSPAPSPGEDICL GLHTEEALCATQACPEGWS
429	1779	A	3464	583	3	DALDRRYLERCHPAAGGWVGEGE*ALCQKT/ RFSGVLEPPLPSLKDGGRFPAWT*RSCSKSLR AAFTSQFFPSRRSRASPGSAP\GNGQNLTEQHP CPGSCDPQVLSASWM*VEHRSKFRPPP*NSTI PPES/RS*QGGTVQTGQHSSGREAGSWRARGR NAGRR*KGGGKIGTKQGAVRARKECRGEMA SGETDSE
430	1780	A	3473	2802	270	FRMRIFLHCPWNQQMWKIWNLLETSLESCKA HLSIQKLLKER\Q\QLPVFKHRDSIVETLKRHR VVVVAGET\GSGKSTQVPHFLLEDLLLNEWE ASKCNIVCTQPRRISAVSLANRVCDELGCENG PGGRNSLCGYQIRMESRACESTRLLYCTTGV LLRKLQEDGLLSNVS/HMFIVDEV\HER\SVQS DFLLIILKEILQKRSDLHLILMSATVDSEKFST YFTHCPILRISGRSYPVEVFHLEDIIEETGFVLE KDSEYCQKFLEEEEEVTINVTSKAGGIKKYQE YIPVQTGAHADLNPFYQKYSSRTQHAILYMN PHKINLDLILELLAYLDKSPQFRNIEGAVLIFL PGLAHIQQLYDLLSNDRRFYSERYKVIALHSI LSTQDQAAAFTLPPPGVRKIVLATNIAETGITI PDVVFVIDTGRTKENKYHESSQMSSLVETFVS KASALQRQGRAGRVRDGFCFRMYTRERFEG FMDYSVPEILRVPLEELCLHIMKCNLGSPEDF LSKALDPPQLQVISNAMNLLRKIGACELNEPK LTPLGQHLAALPVNVKIGKMLIFGAIFGCLDP VATLAAVMTEKSPFTTPIGRKDEADLAKSAL AMADSDHLTIYNAYLGWKKARQEGGYRSEI TYCRRNFLNRTSLLTLEDVKQELIKLVKAAGF SSSTTSTSWEGNRASQTLSFQEIALLKAVLVA GLYDNVGKIIYTKSVDVTEKLACIVETAQGK AQVHPSSVNRDLQTHGWLLYQEKIRYARVY LRETTLITPFPVLLFGGDIEVQHRERLLSIDGW TYFQAPVKIAVIFKQLRVLIDSVLRKKLENPK MSLENDKILQIITELIKTENN
431	1781	A	3474	1	441	FRPAPGHVQP*GGSSAAAGGGLLSHPRPCQQ PCPPAPAPSRPRSLGSLGQRVPAALATAAQEL PATLGGDGGKPALTAGEAALPGLHRSGVPAA AARC*PCT/SRPT*STLSPTQAAWWCRPSRRQ QRGEASTGGASGRRCGSCFQV

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
432	1782	A	3478	416	23	QLRRLTLPNFKTY/YSS*IIEIAWH**KNMQID QWFRRESPEIDLCKYS*LSFDKEAKAIK/WKE CSLFNKWC/YKNWM/LHVQKKRI*VQTLHPS QKLK\SKWIKDLNVECRITKLLDQEYPGDLGY SRALNSGSR
433	1783	A	3504	1876	552	CLAPCSPQPEKNGMQPLLLLLPPLLYQQLLHS SLGAPGESTLLVRTSKLLVGLGLQLLVWLLL QTRSLLALQLHLTSSAPLLAAPTAVCSCSRCS APRSRCVARPAARTGLPTPAPASSPAPAASPA PAASPAPAESTA\PQPLILLPKP/PPAPGAPPPRP GAPPPRPAASPSPAASPAPPAASPVLTASPPLP AASPSPAASPAPPAASPVLTASPPLPAASPSPA ASPAPPAASPVLTASPPLPAASPALAASPVHT ASPPVHVASPPVHTASPPVHV
434	1784	A	3516	142	590	GGVNRPRSETEQVKTPVLISSWDYRHPPPRPA SFFVFLV*TGF\TALARMVLISWPCDLPTSASQ SAGITGVRHHA\RLLYFEQESHSVTQAGW\VQ WHNLGSLQPLSLEDRLSPGVLGCSALCRSGV RTKFGINMVTSRERGTTRLPKEG
435	1785	A	3529	1	3161	MSLVRAALEALDELDLFGVKGGPQSVIHVLA DEVQHCQSILNSLLPRASTSKEVDASLLSVVS FPAFAVEDSQLVELTKQEIITKLQGRYGCCRF LRDGYKTPKEDPNRLYY/ENPAELKLFENIEC EWPLFWTYFILDGVFSGNAEQVQEYKEALEA VLIKGKNGVPLLPELYSVPPDRVDEEYQNPHT VDRVPMGKLPHMWGQSLYILGSLMAEGFLA PGEIDPLNRFSTVPKPDVVVQVYPSLPHGCS SKSPSHQCTIISIRTTRKITAPVSILAETEEIKTIL KDKGIYVETIAEVYPIRVQPARILSHIYSSLEIF LPFLNSVSGCNNRMKLSGRPYRHMGVLGTSK LYDIRKTIFTFTPQFIDQQQFYLALDNKMIVE MLRTDLSYLCSRWRMTGQPTITFPISHSMLDE DGTSLNSSILAALRKMQDGYFGGARVQTGKL SEFLTTSCCTHLSFMDPGPEGKLYSEDYDDN YDYLESGNWMNDYDSTSHARCGDEVARYL DHLLAHTAPHPKLAPTSQKGGLDRFQAAVQT TCDLMSLVTKAKELHVQNVHMYLPTKLFQA SRPSFNLLDSPHPRQENQVPSVRVEIHLPRDQ SGEVDFKALVLQLKETSSLQEQADILYMLYT MKGPDWNTELYNERSATVRELLTELYGKVG EIRHWGLIRYISGILRKKVEALDEACTDLLSH QKHLTVGLPPEPREKTISAPLPYEALTQLIDEA SEGDMSISILTQEIMVYLAMYMRTQPGLFAE MFRLRIGLIIQVMATELAHSLRCSAEEATEGL MNLSPSAMKNLLHHILSGKEFGVERSVRPTD SNVSPAISIHEIGAVGATKTERTGIMQLKSEIK QSPGTSMTPSSGSFPSAYDQQSSKDSRQGW QRRRRLDGALNRVPVGFYQKVWKVLQKCH GLSVEGFVLPSSTTREMTPGEIKFSVHVESVL NRVPQPEYRQLLVEAILVLTMLADIEINHSIGS IIAVEKIVHIANDLFLQEQKTLGADDTMLAKD PASGICTLLYDSAPSGRFGTMTYLSKAAATY VQEFLPHSICAMQ
436	1786	A	3546	73	393	CP*LTWELLEVKKAEVLQDSLDGRYSTPSSCL EQPDSCRPYGRSFYALEEKHVIFSLDVGETDN

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437	1787	A	3554	5157	2939	PXSARSCWMRKG  AVRAEPGLEELSSGLRAHSPSATTVCEPEAQG SASGCRYAAHPHWGLGGAAAAGGSWEPQPP RPVCEPAGRGKPHPPAAPRSPLLPGSRRPHA AQPGARARTSPPPASARNMAARPAATLAWSL LLLSSALLREGCRARFVAERDSEDDGEEPVVF PESPLQSPTVLVAVLARNAAHTLPHFLGCLER LDYPKSRMAIWAATDHNVDNTTEIFREWLK NVQRLYHYVEWRPMDEPESYPDEIGPKHWP TSRFAHVMKLRQAALRTAREKWSDYILFIDV DNFLTNPQTLNLLIAENKTIVAPMLESRGLYS NFWCGITPKGFYKRTPDYVQIREWKRTGCFP VPMVHSTFLIDLRKEASDKLTFYPPHQDYTW TFDDIIVFAFSSRQAGIQMYLCNREHYGYLPIP LKPHQTLQEDIENLHVQIEAMIDRPPMEPSQ YVSVVPKYPDKMGFDEIFMINLKRKGQGGD RWLRTLYEQEIEVKIVEAVDGKALNTSQLKA LNIEMLPGYRDPYSSRPLTRGEIGCFLSHYSV WKEVIDRELEKTLVIEDDVRFEHQFKKKLMK LMDNIDQAQLDWELIYIGRKRMQVKEPEKA
438	1788	A	3563	130	527	VPNVANLVEADYSYWTLGYVISLEGAQKLV GANPFGKMLPVDEFLPVMYNKHPVAEYKEY YESRDLKAFSAEPLLIYPTHYTGQPGYLSDTE TSTIWDNETVATDWDRTHAWKSRKQSRIYSN AKNTEALPPPTSLDTVPSRDEL IFFNSSSLFCRVFCLFLRWSFTLVAQARVQ*C
						NLSSLQPLPPGFK*FSCLSPPRS*DYRRPPPRPA NFLYF**RQGFTVLGQAGLELLT/S/GDPPTSA SQSAGITGVSHRAWPVHAISTHISLVKTRPSLT TLG
439	1789	A	3565	446	1834	LLQPAMRKSPGLSDCLWAWILLLSTLTGRSY GQPSLQDELKDNTTVFTRILDRLLDGYDNRL RPGLGERVTEVKTDIFVTSFGPVSDHDMEYTI DVFFRQSWKDERLKFKGPMTVLRLNNLMAS KIWTPDTFFHNGKKSVAHNMTMPNKLLRITE DGTLLYTMRLTVR\AECPMAFGRDFPM\D\AH ACPLKFGSYAYTRAEVVYEWTREPARSVVV AEDGSRLNQYDLLGQTVDSGIVQSSTGEYVV MTTHFHLKRKIGYFVIQTYLPCIMTVILSQVSF WLNRESVPARTVFGVTTVLTMTTLSISARNSL PKVAYATAMDWFIAVCYAFVFSALIEFATVN YFTKRGYAWDGKSVVPEKPKKVKDPLIKKN NTYAPTATSYTPNLARGDPGLATIAKSATIEP KEVKPETKPPEPKKTFNSVSKIDRLSRIAFPLL FGIFNLVYWATYLNREPQLKAPTPHQ
440	1790	A	3568	1	350	STSSCFPAAAAAIMREIVHLQAGQCGNQIGAK FWEVISDEHGIDPTGTYHGDSDLQLERINVYY NEATGEAPVPSPTALRGPRGPCLG*RPPVPAG GKYVPRAVLVDMEPGTMDSV
441	1791	A	3569	2	1751	FVAVAGAVSGEPLVHWCTQQLRKTFGLDVS EEIIQYVLSIESAEEIREYVTDLLQGNEGKKGQ FIEELITKWQKNDQELISDPLQQCFKKDEILDG QKSGDHLKRGRKKGRNRQEVPAFTEPDTTAE VKTPFDLAKAQENSNSVKKKTKFVNLYTREG QDRLAVLLPGRHPCDCLGQKHKLINNCLICG RIVCEQEGSGPCLFCGTLVCTHEEQDILRGDS NKSQKLLKKLMSGVENSGKVDISTKDLLPH QELRIKSGLEKAIKHKDKLLEFDRTSIRRTQVI

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  DDESDYFASDSNQWLSKLERETLQKREEELR ELRHASRLSKKVTIDFAGRKILEEENSLAEYH SRLDETIQAIANGTLNQPLTKLDRSSEEPLGVL VNPNMYQSPPQWVDHTGAASQKKAFRSSGF GLEFNSFQHQLRIQDQEFQEGFDGGWCLSVH QPWASLLVRGIKRVEGRSWYTPHRGRLWIAA TAKKPSPQEVSELQATYRLLRGKDVEFPNDY PSGCLLGCVDLIDCLSQKQFKEQFPDISQESDS PFVFICKNPQEMVVKFPIKGNPKIWKLDSKIH
442	1792	A	3576	1	2019	QGAKKGLMKQNKAV  MPRSHTGERLCEGKEGSQCAENFSPNLSVTK KTAGVKPYECTICGKAFMRLSSLTRHMRSHT AIRANEKPYKCKEC\GRAFSLSQILSK\HERSH TGEKPYKCKQCGKTFIYHQPFQRHERTHIGEK PYECKQCGKALSCSSSLRVHERIHTGEKPYEC KQCGKAFSCSSSIRVHERTHTGEKPYACK\EC GKAFIS\TTSVLTHMITHNGDRPYKCKECGKA FIFPSFLRVHERIHTGEKPYKCKQCGKAFRWS TSIQIHERIHTGEKPYKCKECGKAFSRISYFRIHERT HTGEKPYECKKCGKTFNYPLDLKIHKRNHTG EKPYECKECAKTFISLENFRRHMITHTGDGPY KCRDCGKVFIFPSALRTHERTHTGEKPYECKQ CGKAFSCSSYIRIHKRTHTGEK\PYECKECGK AFIYPTSFQGHMRMHTGEKPYKCKECGKAFS LHSSFR\RHTRIHNYEKPLEC*Q\CGKAFSVSTS LKKPMRNAQSDRKLY\KCEK*EKVFNSNRCF QSCENSH*REKSCQCK*YRKRDTR*FMYSQV PHNHVSVSNGPYR\CGSPIRLYNT*NISINRNL VAVVTP*CSTLFKCLWCWCKRAALSVV*\IVQ DSGRGRWLTPVIPALWEAKAGGSRGQEIKTIL ANTVKPHLY
443	1793	A	3578	287	114	DFYERKFEQFIEGHKQIVNKWRDLLCSWKRK
444	1794	A	3582	3335	1909	LSIIKKSVLQNNL*FSAASMRFQKVFF HLFFSLFLAAMAMTGSTPCSSMSNHTKERVT MTKVTLENFYSNLIAQHEEREMRQKKLEKV MEEEGLKDEEKRLRRSAHARKETEFLRLKRT RLGLEDFESLKVIGRGAFGEVRLVQKKDTGH VYAMKILRKADMLEKEQVGHIRAERDILVEA DSLWVVKMFYSFQDKLNLYLIMEFLPGGDM MTLLMKKDTLTEEETQFYIAETVLAIDSIHQL GFIHRDIKPDNLLLDSKGHVKLSDFGLCTGLK KAHRTEFYRNLNHSLPSDFTFQNMNSKRKAE TWKRNRRQLAFSTVGTPDYIAPEVFMQTGYN KLCDWWSLGVIMYEMLIGYPPFCSETPQETY KKVMNWKETLTFPPEVPISEKAKDLILRFCCE WEHRIGAPGVEEIKSNSFFEGVDWEHIRERPA AISIEIKSIDDTSNFDEFPESDILKPTVATSNHPE TDYKNKDWVFINYTYKRFEGLTARGAIPSYM KAAK
445	1795	A	3584	1	6169	RTRGIEKRFAYSFLQQLIRYVDEAHQYILEFD GGSRGKGEHFPYEQEIKFFAKVVLPLIDQYFK NHRLYFLSAASRPLCSGGHASNKEKEMVTSL FCKLGVLVRHRISLFGNDATSIVNCLHILGQT LDARTVMKTGLESVKSALRAFLDNAAEDLE KTMENLKQGQFTHTRNQPKGVTQIINYTTVA LLPMLSSLFEHIGQHQFGEDLILEDVQVSCYRI LTSLYALGTSKSIYVERQRSALGECLAAFAGA FPVAFLETHLDKHNIYSIYNTKSSRERAALSLP TNVEDVCPNIPSLEKLMEEIVELAESGIRYTQ

SEQ ID NO: of nucl-eotide sequence	SEQ ID NO: of peptide seq- uence	Method	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, R=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamic, R=Agnine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, y=possible nucleotide deletion, \=possible nucleotide insertion MPHVMEVILPMICSYMSRWWEHGPENNPER AEMCCTALNSEHMNTLLGNILKIIYNNLGIDE GAWMKRLAVFSQPIINKVKPQLLKTHFLPLM EKLKKKAATVVSEEDHLKAEARGDMSEAEL LILDEFTTLARDLYAFYPLLIRFGDYNRAKWL KEPNPEAEELFRMVAEVFIYWSKSHNFKREE QNFVVQNEINNMSFLITDTKSKMSKAAVSDQ ERKKMKRKGGRYSMQTSLIVAALKRLLPIGL NICAPGDQELIALAKNRFSLKDTEDEVRDIRS NIHLQGKLEDPAIRWQMALYKDLPNRTDDTS DPEKTVERVLDIANVLFHLEQKSKRVGRRHY CLVEHPQRSKKAVWHKLLSKQRKRAVVACF RMAPLYNLPRHRAVNLFLQGYEKSWIETEH YFEDKLIEDLAKPGAEPPEDEGTKRVDPLHQ LILLFSRTALTEKCKLEEDFLYMAYADIMAKS CHDEEDDDGEEEVKSFEEKEMEKQKLLYQQ ARLHDRGAAEMYLQTISASKGETGPMVAAT LKLGIAILNGGNSTVQQKMLDYLKEKKDVFFQSLAGILMQSCSVLDLNAFERQNKAEGIGM VTEEGSGEKVLQDDEFTCDLFFFLQLLCEGH NSDFQNYLRTQTGNNTTVNIIISTVDYLLRVQ ESISDFYWYYSGKDVIDEQGQRNFSKAQVA KQVFNTLTEYIQGPCTGNQQSLAHSRLWDAV VGFLHVFAHMQMKLSQDSSQIELLKELMDLQ KDMVVMLLSMLEGNVVNGTIGKQMVDMLV ESSNNVEMILKFDMFLKLKDLTSSDTFKEYD PDGKGVIFKRDFHKAMESHKHYTQSETEFLL SCAETDENFTLDYEEFVKRFHEPAKDIGFNVA VLLTNLSEHMPNIDTRLQTFLELAESVLNYFQP FLGRIEMGSAKRIERVYFEISESSRTQWEKPQ VKESKROFIFDVVNEGGEKEKMELFVNFCED TIFEMQLAAQISESDLNERSANKEESEKERPEE QFRMAFFSILTVRSALFALRYNILTLMRMLS LKSLKQMKKVKKMVKMTVKDMVTAFFSSYWSI FMTLLHFVASVFRGFFRIICSLLLGGSLVEGA KKIKVAELLANMPDPTQDEVRGDGEEGERKP LEAALPSEDLTDLKELTESDLLSDIFGLLKR PMYWDKSPGGIFGRVSSENA KVTSLDSSSNIA KVYSLDSSNIRIAVHYVLEESSGYMEPTVRIL PILHTVISFCIIGYYCLKVPLVIFKREKEVARK LEFDGLYTIFPAGLSDLKGQWDRLVINTQSFP NYWMCKPKREKEEKEEKEETKSEPEKAEGEDGE KEEKAKEDKGKQKLRQUTHTHRYGEPEVPESA FWKLETGGYYKLPHNPNAGLSDLMNPYPMPPVQE KFQEQKAKEEKEEKEEKEETKSEPEKAEGEDGE KEEKAKEDKGKQKLRQUTHTHRYGPPVPESA FWKSTDDSSHRIAVHYVLEESSGYMEPTVRIL PILHTVISFCIIGYYCLKVPLVYTTVVAFNFFRKFYNKSEDGTPDAKCOWEFFPAGCCWEFFPAGCCRUEFFFRAGCECWEFFPAGCCRUEFFFRAGCGCWEFFPAGCCRUEFFFRAGCECWEFFPAGCCRUEFFFRAGCECWEFFPAGCCRUEFFFRAGCEC
446	1796	A	3592	1	355	AGLELLNSDDPPALASQSAGITGVTRTPSLFF* DTVLLCCSGWSAVAPSRLTAALFS*AQAVCL SLPRSWDYRRW/PPHPANFCIFCRDE/SLA/ML PRLVSNSWTQAILLPRPPKMLGLQV

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						GR*V*GAAASQGA/GSLRPRPTGPGHPGAWL QVWGAAAVCAGPAM*/AVRAKRGPRAG*EP NSPWRSGVLAA\RAVGAGPWP*P*PGCS*ARG PSSRSAPGLASGPAAPLLQGVHSSAGPLLCYI NGTLALGLKP**AWGWGEWRPKG
448	1798	A	3604	3115	557	FRRKGGGGPKDFGAGLKYNSRHEKVNGLEE GVEFLPVNNVKKVEKHGPGRWVVLAAVLIG LLLVLLGIGFLVWHLQYRDVRVQKVFNGYM RITNENFVDAYENSNSTEFVSLASKVKDALKL LYSGVPFLGPYHKESAVTAFSEGSVIAYYWSE FSIPQHLVEEAERVMAEERVVMLPPRARSLKS FVVTSVVAFPTDSKTVQRTQDNSCSFGLHAR GVELMRFTTPGFPDSPYPAHARCQWALRGD ADSVLSLTFRSFDLASCDERGRHLV\TVYNT\L SPMEPHA\LVQLCGTYPPSYNLTFHS\S\QNVL LITLITNTERRHPG\FEATFFQLPRMSSCGGRL RKAQGTFNSPYYPGHYPPNIDCTWNIEVPNN QHVKVRFKFFYLLEPGVPAGTCPKDYVEING EKYCGERSQFVVTSNSNKITVRFHSDQSYTDT GFLAEYLSYDSSDPCPGGFTCRTGRCIRKELR CDGWADCTDHSDELNCSCDAGHQFTCKNKF CKPLFWVCDSLNDCGDNSDEQGCSCP\AQTF RCSNGKCLSKSQQCNGKDDCGDGSDEASCP KVNVVTCTKHTYRCLNGLCLSKGNPECDGK EDCSDGSDEKDCDCGLRSFTRQARVVGGTD ADEGEWPWQVSLHALGQGHICGASLISPNWL VSAAHCYIDDRGFRYSDPTQWTAFLGLHDQS QRSAPGVQERRLKRIISHPFFNDFTFDYDIALL ELEKPAEYSSMVRPICLPDASHVFPAGKAIWV TGWGHTQYGGTGALILQKGEIRVINQTTCEN LLPQQITPRMMCVGFLSGGVDSCQGDSGGPL SSVEADGRIFQAGVVSWGDGCAQRNKPGVY TRLPLFRDWIKENTGV
449	1799	A	3618	2	613	FVSGSPWRMDGSTERLEARRPAGRLPWSSRQ EMTRRPSLMAGRQHGWSAQQSATVANPVPG ANPDLLPHFLGEPEDVYIVKNKPVLLVCKAV PATQIFFKCNGEWVRQVDHVIERSTDGSSGLP TMEVRINVSRQQVEKVFGLEEYWCQCVAWS SSGTTKSQKAYIRIAYLRKNFEQEPLAKEVSL EQGIVLPCRPPEGIPPAE
450	1800	A	3620	1	2676	MEPSLGQGMDLTCPFGVSPACGAQASWSIFG ADAAEVPGTRGHSQQEAAMPHIPEDEEPPGE PQAAQSPAGQQGPPTAGVSCSPTPTIVLTGDA TSPEGETDKNLANRVHSPHKRLSHRHLKVST ASLTSVDPAGHIIDLVNDQLPDISISEEDKKKN LALLEEAKLVSERFLTRRGRKSRSSPGDSPSA VSPNLSPSASPTSSRSNSLTVPTPPEGDEADVS SPHPGEPNVPKGLADRKQNDQRKVSQGRLAP RPPPVEKSKEIAIEQKENFDPLQYPETTPKGLA PVTNSSGKMALNSPQPGPVESELGKQLLKTG WEGSPLPRSPTQDAAGVGPPASQGRGPAGEP MGPEAGSKAELPPTVSRPPLLRGLSWDSGPEE PGPRLQKVLAKLPLAEEEKRFAGKAGGKLAK APGLKDFQIQVQPVRMQKLTKLREEHILMRN QNLVGLKLPDLSEAAEQEKGLPSELSPAIEEE ESKSGLDVMPNISDVLLRKLRVHRSLPGSAPP LTEKEVENVFVQLSSAFRNDSYTLESRINQAE RERNLTEENTEKELENFKASITSSASLWHHCE HRETYQKLLEDIAVLHRLAARLSSRAEVVGA

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						AELMEFKKLANQNSSRSCGPSEDGVLRTARS MSLTLGKNMPRRRVSVAVVPKFNALNLPGQ TPSSSSIPSLPALSESPNGKGSLPVTSALPALLE NGKTNGDPDCEASAPALTLSCLEELSQETKA RMEEEAYSKGFQEGLKKTKELQDLKEEEEEQ KSESPEEPEEVEETEEEEKDPRSSKLEELVHFL QVMYPKLCQHWQVIWMMAAVMLVLTVVL GLYNSYNSCAEQADGPLGRSTCSAAQKDSW WSSGLQHEQPTEQ
451	1801	A	3623	504	198	QLIQHQTVHTGRKLYECKECGKAFNQGSTLI RHQRIHTGEKPYECKVCGKAFRVSSQLKQHQ RIHTGERPYQCKELKGRGAEMLAVLAVKEQ NRTPVNYGK
452	1802	A	3628	2	195	MTCLHSAKAFHY*SSCSFSCEEGFALIGPEVV QCTALGVWTAPAPVCIAVQCQHLEALNEGT MG*DYPFTAFAYGSSCKYECHTVYRVRGLD MLHSRGCYLWNGHFTT*EAISCEPLERPCH*S V*CSFSCEEGFALIGPEVVQCTALGVWTAPAP VCIAVQCQHLEALNEGTMG
453	1803	A	3637	662	142	IQAKGLGIWHVPNKSPMQHWR\KGSLLRYRT DTGFLQTLGHNLLGIYQKYPVKYGEGKCWT DNGPVIPVVYDFGDAQKTASYYSPYGQREFT AGFVQFRVFNNERAANALCAGMRVTGCNTE HHCIGGGGYFPEASPQQCGDFSGFDWSGYGT \HVGYSSSREITE\AAVLLFYR
454	1804	A	3641	1	362	TQVHPAMLGLDELGRSGCGHCTQADLRFGD AAGRDPGQDNDRNTAEPAFPPPPRVMAAAA ALRAPAQSSVTFEDVAVNFSLEEWSLLNEAQ GCLYHDVMLETLTLISSLGKVLILNCDLS
455	1805	A	3646	2.	414	AAAGRGASGALTGEGGGEQGRRVGLGSRAH SLLLGPTFNSCQVSSQPPRVAGLGLPLKHEPS RPQPPSPRGPRTVRAGVPGAHPQDTPCPEFVR PRKVPLVGEAPGLPPEERSRGWRRDTPGLQE SRVRAPSYDDIT
456	1806	A	3656	396	8	QIVSFNSYLTLYTKNNLKSMKDLNVNTEMIK LLELKNIHNLG*AKFFLN*IQKALIKRKILIHW P/LIKIK/SFCSLSDTIKKMKRQTIVWEQTFIIHI SVKELVSRIYEAFLQFNKTVNRPVFDIKKEQK F
457	1807	A	3660	14	1961	SEAKLGGPTGMDLWQLLLTLALAGSSDAFSG SEATAAILSRAPWSLQSVNPGLKTNSSKEPKF TKCRSPERETFSCHWTDEVHHGTKNLGPIQLF YTRRNTQEWTQEWKECPDYVSAGENSCYFN SSFTSIWIPYCIKLTSNGGTVDEKCFSVDEIVQ PDPPIALNWTLLNVSLTGIHADIQVRWEAPRN ADIQKGWMVLEYELQYKEVNETKWKMMDP ILTTSVPVYSLKVDKEYEVRVRSKQRNSGNY GEFSEVLYVTLPQMSQFTCEEDFYFPWLLIIIF GIFGLTVMLFVFLFSKQQRIKMLILPPVPVPKI KGIDPDLLKEGKLEEVNTILAIHDSYKPEFHS DDSWVEFIELDIDEPDEKTEESDTDRLLSSDH EKLHINLGVKDGDSGRTSCCEPDILETDFNAH DIHEGTSEVAQPQRLKGEADLLCLDQKNQNN SPYHDACPATQQPSVIQAEKNKPQPLPTEGAE STHQAAHIQLSNPSSLSNIDFYAQVSDITPAGS VVLSPGQKNKAGMSQCDMHPEMVSLCQENF LMDNAYFCEADAKKCIPVAPHIKVESHIQP\S LNQEDIYITTESLT\TAAGSP\GTGEHVPGSEM

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458	1808	A	3663	154	462	SCGYVSTDQLNKIMP TRAPASGRSGAGLALSANAPDSGGHPGATEG
					}	PAGSLAHASGSARGTWRVRGRGSHGWERTV GAGGCANPVPALHSCASAPRGTGRVSALGPK TGSSPLSSPKG
459	1809	A	3664	902	135	LGKYNTSMALFDFVLHNSTGEIRYITEDDVIQ SQNALGKYNTSMALFESNSFEKTILESPYYVD LNQTLFVQVSLHTSDPNLVVFLDTCRASPTSD FASPTYDLIKSGCSRDETCK\VYPLFGHYGRF QFNAFKFLRSMSSVYLQCKVLICDSSDHQSRC \NQGCVSRSKRDISSYKWKTDSIIGPIRLKRDR SA\NGNSGFQHETHAEETPNQPFNSVHLFSFM VLALNVVTVATITVRHFVNQRADYQ\YQKLQ NY
460	1810	A	3670	850	557	LGILMSPQVEAGEI*ALLTPPPGCMQFSPLTL/P K*WVSPGLTP/PPPEVPSVFLVEPGLPHAGQA GLDLL\TSGDPPASTSQSARTTDVSHRAQPLAI S
461	1811	A	3671	2472	2099	IGVLAFETGSCSVTRLYCIGIIMPHCSLDLAGS\ TSAFRIAGTTSVHHHPQLTFFFFWIETGSHCV VQTGL*LLALSNPPALASQIAGISGMSHRAWP GLVLYSLEFSLLCASQSLIMLFTCYNE
462	1812	A	3672	394	110	VKPVNGESKRD*GADTQTCEGEADEQLQT\N CYYD/STKSFFYISCG*K\RKPTWAENRRLNA KMFGIPLHSNSDPWGYEEREVIGFHRSRVSRG HGS
463	1813	A	3673	348	1	QRNPFSAGHPQRPPTSGSQSELLAQPRLRPGR KSSFSRDQDVW*SQAVPKRQ*QRNPFSAGHP QRPPTSGSQSELLAQPRLRPGRKSSFSRDQDV WPGQKPRPSQQQHQMCASPTLGQRSPFALEP VPAYHGGRDPFASARPSPVGIPKPRAAPAGG GWRRIRPKSSTK
464	1814		3676	2253	320	PVIQRCSQPYGFSLLISFFLKCVSETSQQPPSR KVFQLLPSFPTLTRSKSHESQLGNRIDDVSSM RFDLSHGSPQMVRRDIGLSVTHRFSTKSWLS QVCHVCQKSMIFGVKCKHCRLKCHNKCTKE APACRISFLPLTRLRRTESVPSDINNPVDRAAE PHFGTLPKALTKKEHPPAMNHLDSSSNPSSTT FSTPSSPAPFPTSSNPSSATTPP\NPSP\GQR\DSR FNFPSC/AYFIHHR\Q\QFIFPDISAFAHAAPLPE AADGTRLDDQPKADVLEAHEAEAEEPEAGK SEAEDDEDEVDDLPSSRRPWRGPISRKASQTS VYLQEWDIPFEQVELGEPIGQGRWGRVHRGR WHGEVAIRLLEMDGHNQDHLKLFKKEVMN YRQTRHENVVLFMGACMNPPHLAIITSFCKG RTLHSFVRDPKTSLDINKTRQIAQEIIKGMGY LHAKGIVHKDLKSRNVFYDNG\KVVITDFGLF \GISGVVP\EGRRENQLKLSHDWLCYLAPEIVR EMTPGKDEDQLPFSKAADVYAFGTVWYELQ ARDWPLKNQAAEASIWQIGSGEGMKRVLTS VSLGKEVSENLSACWAFDLQERPS\FSLLMD MLEKLPKLNRRLSHPGHF*KSADINSSKVVPR FERFGLGVLESSNPKM
465	1815	A	3679	8	803	IPSPAWWNSTWADTFSLLLALAVALYLGYY WACVLQTHRAFCASNTEDLETVVNHIKHRYP QAPLLAVGISFGGILVLNHLAQARQAAGLVA ALTLSACWDSFETTRSLETPLNSLLFNQPLTA GLCQLVERLSY/E*DLQARTIRQFDERYTSVA

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466	1816	A	3684	3	307	GFLEGLLPWQHWYMSRLLHQYAKAIFQDPE GLPDLRALLPSEDRNS SSQYIVQSKTKIFL*AAREKQ/RHTCRRFSIRLS ANISSQTGEARGQWPSVFKVLKEKKLSTKKS FGQK*GR\RKTFPDKQK/LREFDTTRPTIQEML TGVLQG
467	1817	A	3687	2465	837	ELPTPLIAAHQLYNYVADHASSYHMKPLRMA RPGGPEHNEYALVSAWHISSGSYLDSEGLRHQ DDFDVSLLVCHCAAPFEEQGEAERHVLRLQF FVVLTSQRELFPRLTADMRRFRKPPRLPPEPE APGSSAGSPGEASGLILAFGPAPLFPPLAAEVG MARARLAQLVRLAGGHCRRDTLWKRLFLLE PPGPDRLRLGGRLALAELEELLEAVHAKSIGD IDPQLDCFLSMTVSWYQSLIKVLLSRFPQSCR HFQSPDLGTQYLVVLNQKFTDCFVLVFLDSH LGKTSLTVVFREFFPVQPQDSESPPAQLVSTY HHLESVINTACFTLWTRLL*GSGLDH*MSLFL ESWAYQIACQRQD*PALLGPRASQTLSDTKG FVTMS*GSAAPAWQQEPPSPNTHSH*PIQDSR ESGQPRGPLGPFWGTPFGPPGRVSGVHTGWQ TPPRAPLPESCPLVPLTTVSHLCPLSLRVFTSHL DITAGHSHRDDTWVPIPALPLKHLRPPSSPFA LGPWVSHPLMRWVQKLSHLHSNPGTGFSMG GKQQRN
468	1818	A	3691	960	499	QTCRKDKRAIYPHFQNE*MNEIKAI*SGTGGI QCFHSQNDSAFFFFLFLLETEFCSAA/TVQWH DFLSMQPPPPGFKQFTCLSLLSSWNYRR\PPPF PGNF\*FLVKTGFPHVGQTGFELLTSSDLAPLA SQNGGITGMSPCAWPFFFFFFGLC
	1819	A	3714	4747	495	MAYSWQTDPNPNESHEKQYEHQEFLFVNQP HSSSQVSLGFDQIVDEISGKIPHYESEIDENTFF VPTAPKWDSTGHSLNEAHQISLNEFTSKSREL SWHQVSKAPAIGFSPSVLPKPQNTNKECSWG SPIGKHHGADDSRFSILAPSFTSLDKINLEKEL ENENHNYHIGFESSIPPTNSSFSSDFMPKEENK RSGHVNIVEPSLMLLKGSLQPGMWESTWQK NIESIGCSIQLVEVPQSSNTSLASFCNKVKKIR ERYHAADVNFNSGKIWSTTTAFPYQLFSKTK FNIHIFIDNSTQPLHFMPCANYLVKDLIAEILH FCTNDQLLPKDHILSVWGSEEFLQNDHCLGS HKMFQKDKSVIQLHLQKSREAPGKLSRKHEE DHSQFYLNQLLEFMHIWKVSRQCLLTLIRKY DFHLKYLLKTQENVYNIIEEVKKICSVLGCVE TKQITDAVNELSLILQRKGENFYQSSETSAKG LIEKVTTELSTSIYQLINVYCNSFYADFQPVNV PRCTSYLNPGLPSHLSFTVYAAHNIPETWVHR INFPLEIKSLPRESMLTVKLFGIACATNNANLL AWTCLPLFPKEKSILGSMLFSMTLQSEPPVEM ITPGVWDVSQPSPVTLQIDFPATGWEYMKPD SEENRSNLEEPLKECIKHIARLSQKQTPLLLSE EKKRYLWFYRFYCNNENCSLPLVLGSAPGW DERTVSEMHTILRRWTFSQPLEALGLLTSSFP DQEIRKVAVQQLDNLLNDELLEYLPQLVQAV KFEWNLESPLVQLLHRSLQSIQVAHRLYWL LKNAENEAYFKSWYQKLLAALQFCAGKALN DEFSKEQKLIKILGDIGERVKSASDHQRQEVL KKEIGRLEEFFQDVNTCHLPLNPALCIKGIDH DACSYFTSNALPLKITFINANLMGKNISIIFKA

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	1000					SGLIGPLKENTIKKWFSQHNHLKADYEKALR NFFYSCAGWCVVTFILGVCDRHNDNIMLTKS GHMFHIDFGKFLGHAQTFGGIKRDRAPFIFTS EMICYFITEGGIKNPQHFQDFVELCCRAYNIIR KHSQLLLNLLEMMLYAG\LPELSGI\QDLKY VYNNLRPQDTDLEATSHFTKKIKESLECFPVK LNNLIHTLAQMSAISPAKSTSQTFPQESCLLST TRSIERATILGFSKKSSNLYLIQVTHSNNETSL TEKSFEQFSKLHSQLQKQFASLTLPEFPHWW HLPFTNSDHRRFRDLNHYMEQILNVSHEVTN SDCVLSFFLSEAGQQTVEESSPVYLGEKFPDK KPKVQLVISYEDVKLTILVKHMKNIHLPDGSA PSAHVEFYLLPYPSEVRRKTKSVPKCTDPTY NEIVVYDEVTELQGHVLMLIVKSKTVFVGAI NIRLCSVPLDKEKWYPLGNSII*PLLLFSSFGM KSLEKDEFVGGMILSNPIW
470	1820	A	3718	430	75	SHGSISILNLHQGCVFLPSLPAQGLRCYRCLA VLEGASCSVVSCPFLDGVCVSQKVSV/CWQ*/ CPWGARAEGRLSAVVDSQISCCKGDLCNAV VLAAGSPWALCVQLLLSLGSVFLWALL
471	1821	A	3723	891	494	LRQSL/NSVPQAGVQWRDSSLQAPPPRFTPLS CLSLPSSWDYRRLPPCLANFLYF**RRGFTML ARMVLIS*PRDPPASASQ\STEITGGSHRAQHP TDSRDHSERSVKKSHEVISELRMKVIKCKVAF SKNPI
472	1822	A	3734	443	251	GFIET*NFCVSKDTSKKLS/RLPTKWKNVFAN *ISDKGLVSRICQELLRHLDAEQVSSTAGLSL
473	1823	A	3746	3	500	THASGGARSGAGWAGRGVRAGTEAGRGGIF LTLSILRTRDLPSGAMSEGVDLIDIYADEEFNQ DPEFNNTDQIDLYDDVLTATSQPSDDRSSSTE PPPPVRQEPSPKPNNKTPAILYTYSGLRNRRA AVYVGSFSWWTTDQQLIQVIRSIGVYDVGEV KFAENRAK
474	1824	A	3753		. 5262	RPLFAREGGIYAVLVCMQEYKTSVLVQQAG LAALKMLAVASSSEIPTFVTGRDSIHSLFDAQ MTREIFASIDSATRPGSESLLLTVPAAVILMLN TEGCSSAARNGLLLINLLCNHHTLGDQIITQ ELRDTLFRHSGIAPRTEPMPTTRTILMMLLNR YSEPPGSP\ERAALETPIIQQDGSPELLIRSLV GGPSAELLLDLERVLCREGSPGGAVRPLLKRL QQETQPFLLLRTLDAPGPNKTLLLSVLRVIT RLLDFPEAMVLPWHEVLEPCLNCLSGPSSDSE IVQELTCFLHRLASMHKDYAVVLCCLGAKEI LSKVLDKHSAQLLLGCELRDLVTECEKYAQL YSNLTSSILAGCIQMVLGQIEDHRRTHQPINIP FFDVFLRHLCQGSSVEVKEDKCWEKVEVSSN PHRASKLTDHNPKTYWESNGSTGSHYITLHM HRGVLVRQLTLLVASEDSSYMPARVVVFGG DSTSCIGTELNTVNVMPSASRVILLENLNRFW PIIQIRIKRCQQGGIDTRVRGVEVLGPKPTFWP LFREQLCRRTCLFYTIRAQAWSRDIAEDHRRL LQLCPRLNRVLRHEQNFADRFLPDDEAAQAL GKTCWEALVSPLVQNITSPDAEGVSALGWLL DQYLEQRETSRNPLSRAASFASRVRLCHLL VHVEPPPGPSPEPSTRPFSKNSKGRDRSPAPSP VLPSSSLRNITQCWLSVVQEQVSRFLAAAWR APDFVPRYCKLYEHLQRAGSELFGPRAAFML

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						ALRSGFSGALLQQSFLTAAHMSEQFARYIDQ QIQGGLIGGAPGVEMLGQLQRHLEPIMVLSG LELATTFEHFYQHYMADRLLSFGSSWLEGAV LEQIGLCFPNRLPQLMLQSLSTSEELQRQFHLF QLQRLDKLFLEQEDEEKRL*EEEEEEEEEE EKELFIEDPSPAISILVLSPRCWPVSPLCYLYHP RKCLPTEFCDALDRFSSFYSQSQNHPVLDMG PHRRLQWTWLGRAELQFGKQILHVSTVQMW LLLKFNQTEEVSVETLLKDSDLSPELLLQALV PLTSGNGPLTHHEGQDFPHGGVLRLHEPGPQ RSGEALWLIPPQAYLNVEKDEGRTLEQKRNL LSCLLVRILKAHGEKGLHIDQLVCLVLEAWQ KGPNPPGTLGHTVAGGVACTSTDVLSCILHLL GQGYVKRRDDRPQILMYAAPEPMGPCRGQA DVPFCGSQSETSKPSPEAVATLASLQLPAGRT MSPQEVEGLMKQTVRQVQETLNLEPDVAQH LLAHSHWGAEQLLQSYSEDPEPLLLAAGLCV HQAQAVPVRPDHCPVCVSPLGCDDDLPSLCC MHYCCKSCWNEYLTTRIEQNLVLNCTCPIAD CPAQPTGAFIRAIVSSPEVISKYEKALLRGYVE SCSNLTWCTNPQGCDRILCRQGLGCGTTCSK CGWASCFNCSFPEAHYPASCGHMSQWVDDG GYYDGMSVEAQSKHLAKLISKRCPSCQAPIE KNEGCLHMTCAKCNHGFCWRCLKSWKPNH KDYYNCSAMVSKAARQEKFFQDYNERCTFH HQAREFAVNLRNRVSAIHEVPPPRSFTFLNDA CQGLEQARKVLAYACVYSFYSQDAEYMDVV EQQTENLELHTNALQILLEETLLRCRDLASSL RLLRADCLSTGMELLRIQERLLAILQHSAQD FRVGLQSPSVEAWEAKGPNMPGSQPQASSGP EAEEEEEDDDDDVPEWQQDEFDEELDNDSFS YDESENLDQETFFFGDEEEDEDEAYD
475	1825	A	3754	1093	96	GTSRNQHSPKTHA*RSS/WPQPPPLFLPPLQPQ ATGRRRRTRTQQRTAALLTDGTTKTGAAW SRRPSLCWPSRTTGAPGAK*AVLVRSATPTTN PPNPQSPTGAAGKLRAPGNRAG/SEPSSQEPPP DGTR\RPASITGVAQSPATRATPSLPCLHVPAP SRGQTLGVRTTGRASRLTVDRSRLSWPGRSA RSGGGRWRPNAPRGRWPRAP*SWEPGSWTE PWRWPFPAAESPPHRCIYCTNHVSPAGPARPS HVYIIRATINSISHPLCRAQSSPWEAAGVWRR PAQPAPTSDVNINLLRKPRVKRHDLIYQFLGN TLWEEGRQRPPETLQPAR
476	1826	A	3758	901	521	FFFGNGVSPCPQAGV*WHDLDSLQNLPPGFK RFSYLSLPSSW\DYRHVPPRQANFCIF/M*RRG FTMLARMVSIS*PRDLPALASQSAGITGVSHH APPQMDFTFALLCFAPKGCLPRQKEGGTLNLI
477	1827	A	3761	843	575	GVISAHCNLRL/CHLPGSSNSPASASQVAGTIG ARTTPS*IFVFLVETGFHHVSQDGLDLL/NFVI RPRRPLKVLGLQACTRARLPSPLKEL
478	1828	A	3763	267	1240	HILISPHLWSASLDCLEQLSQERHVKGMLLGP PPVNESTKPSPSPWKLTPPMCSIPPVFPPKSGS PTTSWS/PSGHSKLEVERAQTGPFCLHIYCP*P GYTDNTTSLLHYIPFPRL\SGLVCFPAH*FPSY WTGHSFASQAWLRQVPEVSKHLQCPSAESLL TMEYHQPEDPAPGKAGTAEAVIPENHEVLAG PDEHPQDTDARDADGEAREREP/RRPSFAA*P VWGQP\ESPLPEASSAPPGPTLGTLPEVETIRA CSMPQELP*SPRTRQPEPDFYCVKWIPWKGE QTPIITQSTNGPLPSPCHHEHPLSSVEGEAPPA

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479	1829	A	3766	2	2152	YSPIRLLEVCVPLPKIFIKRQAPLKVSLLQDLK DFFQKVSQVYVAIDERLASLKTDTFSKTREEK MEDIFAQKEMEEGEFKNWIEKMQARLMSSS VDTPQQLQSVFESLIAKKQSLCEVLQAWNNR LQDLFQQEKGRKRPSVPPSPGRLRQGEESKIS AMDASPRNISPGLQNGEKEDRFLTTLSSQSST SSTHLQLPTPPEVMSEQSVGGPPELDTASSSE DVFDGHILLGSTDSQVKEKSTMKAIFANLLPG NSYNPIPFPFDPDKHYLMYEHERVPIAVCEKE PSSIIAFALSCKEYRNALEELSKATQWNSAEE GLPTNSTSDSRPKSSSPIRLPEMSGGQTNRTTE TEPQPTKKASGMLSFFRGTAGKSPDLSSQKRE TLRGADSAYYQVGQTGKEGTENQGVEPQDE VDGGDTQKKQLINPHVELQFSDANAKFYCRL YYAGEFHKMREVILDSSEEDFIRSLSHSSPWQ ARGGKSGAAFYATEDDRFILKQMPRLEVQSF LDFAPHYFNYITNAVQQKRPTALAKILGVYRI GYKNSQNNTEKKLDLLVMENLFYGRKMAQ VFDLKGSLRNRNVKTDTGKESCDVVLLDENL LKMVRDNPLYIRSHSKAVLRTSIHSDSHFLSS HLIIDYSLLVGRDDTSNELVVGIIDYIRTFTWD KKLEMVVKSTGILGGQG*MPTVVSPELYRTR FCEAMDNYFLMVPDHCTGLGLNC
480	1830	A	3777	251	3	QGCGSAGTLIHY**ECKMVQLLWKTV*QFLI KLNI\KDPAITLDVYPNEVKNYVRTKTYTQMF I/ANFIMAKSWKQPTHPSVRT
481	1831	A	3779	333	3	EAAIRQPEPNILDVNQIFKDLAMIIHDQGDLID SIEANAESSEVLVERAPGQLQRPA\YYQKKSR KKMCLVVLVQTAIILICERIM*VVYTTKWSPPI VLPVSCFQGQKFN
482	1832	A	3780	2	371	TGGRQGKNDHTSITEKPSRDFNRHLITQNI*M PNQDMKSSSNSLIIRKVQIKPTILYHHIFTRKA KMKTTDKTKYR*GFKAITTLIHCSQDCKLQ*S /L*ENHFMIFPKAEQHITYDTTIPFLR
483	1833	A	3787	43	448	LMKDLSPYVMETHYILNRLNER/RSMWRHIIG KLPNTKDQEKILKAIRGRREVIQGS/RQQYRR PAAFSAAEKARRLWCS/VFNIERRNL/CEYPTK LSFNIKGEMTFSDKTEFTTNRPSLKMLLKDRI QEEGKMF*KEKCFKRKE
484	1834	A	3798	1	727	FFFFETESRSVAQAGVQWCNLGSLQALPPGF\ SHSPASASRVAGTTGTRH*ARLIFYIFSRDGVS PC*PGWS*SPDLVIRPP\RLPKCWDYRREPPRP A*FFVFLVE\QGFTMLARMVSIS*PQ/CDLPAS VSQNAGITGVSHCAWPCLHFCFFGFFFEMESC SVAQAEVQWHDLRSLQAPPPGFTPFSCLSLPG SWDYRRPPPRPANF\CIFSRDGVSPC*PGWSRS PDLVIRPPRPPKVLGLQA
485	1835	A	3802	1	239	FFFFEMECLTVSQAGVQWYNLHSLQPLPPGF KQFSC\LSLPSSWD*RVPTSRPAKF/CVIF*DGV SHCQPGWSAVVQPPLH
486	1836	A	3811	378	98	RYD*SSQSENIP\QKEFLLKYP*CTATLGMRN MSIMKKKSIFSAEFYKVSLPSLLL\HLLAIEWG FHIEIQLTIHQHFLNYELESDFVHIVEYM
487	1837	A	3814	771	320	FDPDWTRAAGIRHEKKPKALAYRRENSPGDL PPPPLPPPEEEASWAL/GAEGSRQHVLPGAGA QWGEESGPGRAPGSPAGAPPR*RGLAP\NSRP SFLSRGQGTSTCSTAGSNSSRGSSSSRGSRGPG RSRSRSQSRSQSQRPGQKRREEPR

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488	1838	A	3818	1	781	FRACLLELIPYAPTLSWTACPPAMAGPRGLLP LCLLAFCLAGFSFVRGQVLFKGCDVKTTFVT HVPCTSCAAIKKQTCPSGWLRELPDQITQDCR YEVQLGGSMVSMSGCRRKCRKQVVQKACCP GYWGSRCHECPGGAETPCNGHGTCLDGMDR NGTCVCQENFRGSACQECQDPNRFGPDCQSV CSCVHGVCNHGPRGDGSCLCFAGYTGPHCD QELPVWQELGFPQNNPRLRKAPNCKCLPG*H RNGLIATPNPCRP
489	1839	A	3822	934	669	FFFSEMESRSVTRLECSGAISAHLRLLGSSNSP ASAS*VAGTIGACHHAQLIFVFLVETGFHHVG QDGLDLL/NLMIHPPRPPKVLGFQA
490	1840	A	3825	79	9748	GCQSCWPAWPRLRRRGPASAGARLGRKAPW GLPGRVQDGRPLRFCFYLRPRAPFIAPVLSGA ASRPEASGDCRAGRETAMATLEKLMKAFESL KSFQQQQQQQQQQQQQQQQQQQQQQPPPP PPPPPPPQLPQPPPQAPPLPPPPPPPPP GPAVAEEPLHRPKKELSATKKDRVNHCLTIC ENIVAQSVRNSPEFQKLLGIAMELFILCSDDA ESDVRMVADECLNKVIKALMDSNLPRLQLEL YKEIKKNGAPRSLRAALWRFAELAHLVRPQK CRPYLVNLLPCLTRTSKRPEESVQETLAAAVP KIMASFGNFANDNEIKVLLKAFIANLKSSSPTI RRTAAGSAVSICQHSRRTQYFYSWLLNVLLG LLVPVEDEHSTLLILGVLLTLRYLVPLLQQQV KDTSLKGSFGVTRKEMEVSPSAEQLVQVYEL TLHHTQHQDHNVVTGALELLQQLFRTPPPEL LQTLTAVGGIGQLTAAKEESGGRSRSGSIVELI AGGGSSCSPVLSRKQKGKVLLGEEEALEDDS ESRSDVSSSALTASVKDEISGELAASSGVSTPG SAGHDIITEQPRSQHTLQADSVDLASCDLTSS ATDGDEEDILSHSSSQVSAVPSDPAMDLNDG TQASSPISDSSQTTTEGPDSAVTPSDSSEIVLD GTDNQYLGLQIGQPQDEDEEATGILPDEASEA FRNSSMALQQAHLLKNMSHCRQPSDSSVDKF VLRDEATEPGDQENKPCRIKGDIGQSTDDDS APLVHCVRLLSASFLLTGGKNVLVPDRDVRV SVKALALSCVGAAVALHPESFFSKLYKVPLD TTEYPEEQYVSDILNYIDHGDPQVRGATAILC GTLICSILSRSRFHVGDWMGTIRTLTGNTFSL ADCIPLLRKTLKDESSYTCKLACTAVRNCVM SLCSSSYSELGLQLIIDVLTLRNSSYWLVRTEL LETLAEIDFRLVSFLEAKAENLHRGAHHYTGL LKLQERVLNNVVIHLLGDEDPRVRHVAAASL IRLVPKLFYKCDQGQADPVVAVARDQSSVYL KLLMHETQPPSHFSVSTITRIYRGYNLLPSITD VTMENNLSRVIAAVSHELITSTTRALTFGCCE ALCLLSTAFPVCIWSLGWHCGVPPLSASDESR KSCTVGMATMILTLLSSAWFPLDLSAHQDAL ILAGNLLAASAPKSLRSSWASEEEANPAATK QEEVWPALGDRALVPMVEQLFSHLLKVINIC AHVLDDVAPGPAIKAALPSI.TNPPSLSPIRRK GKEKEPGEQASVPLSPKKGSEASAASRQSDTS GPVTTSKSSSLGSFYHLPSYLKLHDVLKATHA NYKVTLDLQNSTEKFGGFLRSALDVLSQILEL ATLQDIGKCVEEILGYLKSCFSREPMMATVC VQQLLKTLFGTNLASGFDGLSSNPSKSQGRA QRLGSSSVRPGLYHYCFMAPYTHFTQALADA SLRNMVQAEQENDTSGWFDVLQKVSTQLKT NLTSVTKNRADKNAIHNHIRLFEPLVIKALKQ

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10.						TWSLSCFFVSASTSPWVAAILPHVISRMGKLE QVDVNLFCLVATDFYRHQIEEELDRRAFQSV LEVVAAPGSPYHRLLTCLRNVHKVTTC
491	1841	A	3826	469	302	SNPPASASRVAGITGVHQHAWLIFVFLVEMEF HHVGQAVLKLLISGDLPVSASQSA
492	1842	À	3836	392	88	VAPSPMIMPDLYFYRDPEEIEKEE*AAAEK\EE
						FQSEWTAVV/P/EFTATQSEVADWFKDMQVP SVPIQQFPTEDWST*PIMNDWSATSTAQTTE WVRITTEWP

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493	1843	A	3838	19	380	TPSDMNRAFETDTQSIGEKNRSPSEPDYFERK KFKRS*EKAHIRYKIDQPEDIPLK\EFLCKHSK CTATLSMRNMSLMKKKCSFSEEF\LAFFPSLL VCHLLAIKLGFYIEIHLTTFNNTF
494	1844	A	3845	2	352	FFFLRRSL/DSVAQAEAQWL\ELGLLQAPPPGF KPISLP\GLPSSWDYGRPPPCPANFCIF/M*RRG FTVLARMVLIS*PCDPPTLASQGTAITGMSYH ARPQDIDFLYAHQGRCWFRLL
495	1845		3847	1774	40	DIFFRAKEGMGQDEAQFSVEMPLTGKAYL WADKYRPKKPRFFNRVHTGFEWNKYNQTHY DFDNPPPKIVQGYKFNIFYPDLIDKRSTPEYFL EACADNKDFAILRFHAGPPYEDIAFKIVNREW EYSHRHGFRCQFANGIFQLWFHFKRYRYRR* RPWGTAGRCPRGHSKGASVKLVVTPGPLSGL QGRGFTSHLRPHLSFARPQFPPI*KGGHH*AC HGELRRHWDRLA*GPDATEGALGASFEHEG GQQPPADLTVQADTLHRPSARLGGAHRACPK RRPHRVLWRWARGAWAWRCQAREKQETQG QPCHITGHPLGREAEPAAAGAAPALAHRPPF ARTGSTE\PGPCWRPIRHCRRDPLWTPTLC\RD WPPTHPVLAGGVHFPAAG/IGGCVEVPVSVN VMGTKSH*AVLPPPSTGPGQGGLPEGWGLE KGEGLPPGIPPPGLLTGPW\SMRPVTPSFAHIR TVAPSHSPFSGQEGRGPHGCHSPGR\SGP\AGR LVLQHPTGTSPTEAKRKVPPGPPEGHPTSPVT SPRPPTAPPRHPASSGNSSVCFSKKTCRWEKK SFVLMELAYWQDRMFF
496	1846	A	3849	830	442	AKSPLPLG*IQWR/NLGSLKLRLPGFK*FTCLG LLSSWDYRSLPPRPVNFCILVELGFHHVDQAG LKLLTSSALPALASQSAEITGMSHRIWPLPLLR RPPVIRIRAPPQRLPFNLITSLKALSPNMATF
497	1847	A	3859	2	393	ALRKTRRDGIARTGAQPAASWKGTNNYPWR LEMAGRPGSQEQSKDRGTGSLPPPSQRPLGPS PEGAGPSPPPGIPRGGGSSSSEGP/PQLLFVPR RFPAPKKGLPSDTPHSKAPPTPHLILGGEDSQ VPIL
498	1848	A	3860	253	634	KNASTVYSSQGDPKSFFFLLRWSLALVAQAG EQ*RDLSSLQPPPPGFK*FSCLSLPSSWD\YRCP LPCLANF\*FLVETGFHHVGQADLKLLTSGDP PTSASESAGITGVSHRAWPRIHFLYWKTFFL
499	1849		3863	423	263	APSQISVAFLYAA/DKLFEKEI*KKIPFIIAS/DKI KIGINLTKEVKYLYTENYITLMKEIK/DTDKW KDILY*WIGKINI*KMSTPPKAIYRFNAIPTKIP MTFFTEIEKSIIKFIWNHKKPPNTQSNIEQKE*S FCSILLWVFGGFLWFHMNFMIDFSISVKNVIGI LVGIALNL
500	1850	A	3865	2	15246	LPRGCLWCLQRSPTPARPQPSRPARSPLPLFP DLRPWASDLDIMGDAEGEDEVQFLRTDDEV VLQCSATVLKEQLKLCLAAEGFGNRLCFLEP TSNAQNVPPDLAICCFVLEQSLSVRALQEML ANTVEAGVESSQGGHRTLLYGHAILLRHAH SRMYLSCLTTSRSMTDKLAFDVGLQEDATGE ACWWTMHPASKQRSEGEKVRVGDDIILVSVS SERYLHLSTASGELQVDASFMQTLWNMNPIC SRCEEGFVTGGHVLRLFHGHMDECLTISPADS DDQRRLVYYEGGAVCTHARSLWRLEPLRIS WSGSHLRWGQPLRVRHVTTGQYLALTEDQG LVVVDASKAHTKATSFCFRISKEKLDVAPKR DVEGMGPPEIKYGESLCFVQHVASGLWLTYA

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HSSLKCSNCYMVWGGDFVSPGQQGRISHTDL VIGCLVDLATGLMTFTANGKESNTFFQVEPN TKLFPAVFVLPTHQNVIQFELGKQKNIMPLSA AMFQSERKNPAPQCPPRLEMQMLMPVSWSR MPNHFLQVETRRAGERLGWAVQCQEPLTMM ALHIPEENRCMDILELSERLDLQRFHSHTLRL YRAVCALGNNRVAHALCSHVDQAQLLHALE DAHLPGPLRAGYYDLLISIHLESACRSRSML SEYIVPLTPETRAITLFPPGRSTENGHPRHGLP GVGVTTSLRPPHHFSPPCFVAALPAAGAAEAP ARLSPAIPLEALRDKALRMLGEAVRDGGQHA RDPVGASVEFQFVPVLKLVSTLLVMGIFGDE DVKQILKMIEPEVFTEEEEEDEEEEGEEDEE EKEEDEETAQEKEDEEKEEEAAAEGKEEG LEEGLLQMKLPESVKLQMCHILEYFCDQELQ HRVESLAAFAERYVDKLQANQRSRYGLIIKA FSMTAAETARRTREFRSPPQEQINMLLQFKDG TDEEDCPLPEEIRQDLLDFHQDLLAHCGIQLD GEEEEPEEETTLGSRLMSLLEKVRLVKKKEEK PEEERSAEESKPRSLQELVSHMVVRWAQEDF VQSPLVRAMFSLLHRQYDGLGELLRALPRA YTISPSSVEDTMSLLECLGQIRSLLIVQMGPQE ENLMIQSIGNIMNNKVFYQHPNLMRALGMHE							
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TVMEVMVNVLGGGESKEIRFPKMVTSCCRFL	1						ENLMIQSIGNIMNNKVFYQHPNLMRALGMHE
							TVMEVMVNVLGGGESKEIRFPKMVTSCCRFL

sequence uence uence 914 correspondi ng to first amino acid residue of peptide sequence sequence sequence uence 914 correspondi ng to first amino acid residue of peptide sequence sequ	s, S=Serine, V=Tryptophan, , *=Stop codon,
CYFCRISRQNQRSMFDH QGSTPLDVAAASVIDNN VSYLAGCGLQSCPMLV. ERYLDFLRFAVFVNGES KPECFGPALRGEGGSGL DGPGIRRDRRREHFGEE	NELALALQEQDLEKV AKGYPDIGWKPCGG SVEENANVVVRLLIR LLAAIEEAIRISEDPAR
FYAALIDLLGRCAPEMF LRSLVPLEDLVGIISLPLG MSASFVPDHKASMVLFI VLDVGFLPDMRAAASLI NRYLCLAVLPLITKCAP	HLIQAGKGEALRIRAI QIPTLGKDGALVQPK LDRVYGIENQDFLLH DTATFSTTEMALAV
MLHTVYRLSRGRSLTKA YIRPSMLQHLLRRLVFD TNHYERCWKYYCLPTG TRKLFWGIFDSLAHKKY AIAGALPPDYVDASYSS	DVPILNEFAKMPLKLL GWANFGVTSEEELHL YDPELYRMAMPCLC GKAEKKATVDAEGNF
DPRPVETLNVIIPEKLDS DKIQNNWSYGENIDEEL KDKEIYRWPIKESLKAM EEKTEKKKTAKISQSAQ SAVTLSRELQAMAEQLA ELEAKGGGTHPLLVPYI	LKTHPMLRPYKTFSE MAWEWTIEKAREGE OTYDPREGYNPQPPDL AENYHNTWGRKKKQ
QELLKFLQMNGYAVTR FAFGFLQQLLRWMDISQ EKSPHEQEIKFFAKILLP TPAKVLGSGGHASNKEI RHRVSLFGTDAPAVVNO	GLKDMELDSSSIEKR QEFIAHLEAVVSSGRV LINQYFTNHCLYFLS KEMITSLFCKLAALV CLHILARSLDARTVM
KSGPEIVKAGLRSFFESA KVSQARTQVKGVGQNL QHIAQHQFGDDVILDDV TTKNTYVEKLRPALGEO PQLNEYNACSVYTTKSP CPDIPVLERLMADIGGL	TYTTVALLPVLTTLF VQVSCYRTLCSIYSLG CLARLAAAMPVAFLE PRERAILGLPNSVEEM
ITLPMLCSYLPRWWERG CTAVTSDHLNSLLGNILI KRLAVFAQPIVSRARPEI AGKVVSEEEQLALEAKA VLCRDLYALYPLLIRYV	GPEAPPSALPAGAPPP RIIVNNLGIDEASWM LLQSHFIPTIGRLRKR AEAQEGELLVRDEFS
AEELFRMVGEIFIYWSK: NEINNMSFLTADNKSKN RTKKKRRGDRYSVQTSI MCAPTDQDLITLAKTRY NNLHLQGKVEGSPSLRV	MAKAGDIQSGGSDQE LIVATLKKMLPIGLN YALKDTDEEVREFLH WQMALYRGVPGREE
DADDPEKIVRRVQEVSA KKAVWHKLLSKQRRRA THRACNMFLESYKAAW LSKAGEQEEEEEEVEEK ALTEKSKLDEDYLYMA	AVVACFRMTPLYNLP VILTEDHSFEDRMIDD KRPDPLHQLVLHFSRT YADIMAKSCHLEEG
GENGEAEEEVEVSFEEK HTRGAAEMVLQMISACI GISILNGGNAEVQQKML IQALMQTCSVLDLNAFE DGTVINRQNGEKVMAD EGHNNDFQNYLRTQTGI	KGETGAMVSSTLKL LDYLKDKKEVGFFQS ERQNKAEGLGMVNE DDEFTQDLFRFLQLLC
RLQESISDFYWYYSGKD SVAKQVFNSLTEYIQGP DAVVGFLHVFAHMMMI LDLQKDMVVMLLSLLE DMLVESSSNVEMILKFFI QDYVTDPRGLISKKDFQ	DVIEEQGKRNFSKAM CTGNQQSLAHSRLW KLAQDSSQIELLKEL GNVVNGMIARQMV DMFLKLKDIVGSEAF

eotide sequence USSN   location   corresponding to last amino acid   residue of peptide   sequence   length   location   corresponding to last amino acid   membership   membership   length   l	=Proline, rine,
uence  914  ng to first amino acid residue of peptide residue of peptide sequence  914  ng to first amino acid residue of peptide sequence  92-Glutamine, R=Arginine, S=Se T=Threonine, V=Valine, W=Tryp Y=Tyrosine, X=Unknown, *=Sto /=possible nucleotide deletion, \= nucleotide insertion QFLLSCSEADENEMINCEEFA	rine,
amino acid residue of peptide residue of peptide sequence y=Tyrosine, X=Unknown, *=Sto /=possible nucleotide deletion, \=  nucleotide insertion QFLLSCSEADENEMINCEEFA	
residue of peptide sequence Y=Tyrosine, X=Unknown, *=Sto /=possible nucleotide deletion, \= nucleotide insertion QFLLSCSEADENEMINCEEFA	ntonhan
peptide /=possible nucleotide deletion, \=  sequence nucleotide insertion QFLLSCSEADENEMINCEEFA	
sequence nucleotide insertion QFLLSCSEADENEMINCEEFA	
QFLLSCSEADENEMINCEEFA*	possible
	NRFOEPARDIG
YFRPYLGRIEIMGASRRIERIY	FEISETNRAQW
EMPQVKESKRQFIFDVVNEGO	
FCEDTIFEMQIAAQISEPEGEPI	
AEAGAEGAAGLEGTAA	
VAAAGRALRGLSYRSLRRRV	
ATAVAALLWAAVTRAGAAG	
LWGSLFGGGLVEGAKKVTVT SDEVHGEQPAGPGGDADGEG	
GAGDEEAVHEAGPGGADGA	
PEGAGGLGDMGDTTPAEPPTI	
VDGVEEELPPEPEPEPELEP	
EEVPEPTPEPPKKQAPPSPPPK	KEEAGGEFWG
GGSSGWGLGAGEEAEGDEDE	
TGYMEPALRCLSLLHTLVAFI	
LVIFKREKELARKLEFDGLYIT  OWDRLVLNTPSFPSNYWDKF	
DIYGRERIAELLGMDLATLEIT	
PGLLTWLMSIDVKYQIWKFG	
WYMVMSLLGHYNNFFFAAH	
RTILSSVTHNGKQLVMTVGLI	
AFNFFRKFYNKSÈDEDEPDMI	
FHMYVGVRAGGGIGDEIEDPA	
FDITFFFFVIVILLAIIQGLIIDAI	
KEDMETKCFICGIGSDYFDTT	
HNLANYMFFLMYLINKDETE MYQERCWDFFPAGDCFRKQY	
501 1851 A 3869 467 665 VIVAIYCQLIFDKGAKTIQ*PFG	
LGPCFTPCGKINSEWIRELSVR	
N	
502 1852 A 3888 1042 724 SGMQWRDLTPLQPLPPRFKQI	FSCLSLPGSWD
YRHAP\PLLTNF\*FLVEMGFC	
ASSDQSALASQSAGITGISTAP	GPPFFFLNFEA
GSCSVAQAGVQ	OT TANGOT - SS
503 1853 A 3891 1773 1193 EVDSQSGVQ*QAPGSLQLQTP	
QDYRSSLPHLASCCYYYYYYY VQGGLKLLPSSNPFASAP*TAG	
VQGGLALLPSSNFFASAF*TAG HFNF*MFRKISCIRE*F*HTRIY	
WVLLCYPGWPQIPGLKPSSCL	
APPCPASFFIFHVDRVSPPCPG	
L	
504         1854         B         3896         279         70         MVSKSKSILMSYNHVELTFSD	MKKMPEAFRR
TQKHTIYLIPYQVIFWSTGKDA	AMRSFMMPFY
QKEYYENQ*	
505 1855 A 3899 2 1396 EPGVPTKKTWFDKPDFNRTNS	
NENTKLELRK VPPELNNISKLI	
NLQVAYNGDPEGALIQFATYI	
VLNNRFIKVYWHREGSTQQL0 VQQPILPVVKQSVKERLGPVP	
ASSDLPQVLST\LLA*QKQCIIQ	
LLVSTSAVDNNEAQKKKQEA	
KQEILEKHIETQKMLISKLEKN	
IMKTLEVLTKNITKLKDEVKA	
KTKTQMQKELLDTELDLYKK	
RRKYTELQLEAAKRGILSSGR	GRGIHSRGRGA
VHGRGRGRGRGVPGHAV	VDHRPRALEIS

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  AFTESDREDLLPHFAQYGEIEDCQIDDSSLHA
						VITFKTRAEAEAAAVHGARFKGQDLKLAWN KPVTNISAVETEEVEPDEEEQREIIIA
506	1856	A	3911	1952	919	DAELSGTLSLVLTQCCKRIKDTVQKLASDHK DIHSSVSRVGKAIDKNFDSDISSVGIDGCWQA DSQRLLNEVMVEHFFRQGMLDVAEELCQES GLSVDPSQKEPFVELNRILEALKVRVLRPALE WAVSNREMLIAQNSSLEFKLHRLYFISLLMG GTTNQREALQYAKNFQPFALNHQKDIQVLM GSLVYLRQGIENSPYVHLLDANQWADICDIFT RDACALLGLSVESPLSVSFSAGCVALPALINIK AVIEQRQCTGVWNQKDELPIEVDLG*KSAGY HSIFACPILRQQTTDNNPPMKLVCGHIISRDAL NKMFNGSKLKCPYCPMEQSPGDAKQIFF
507	1857	A	3936	439	18	SHPFSPAPGICPDAPPPLPRPSKGLGHPGTAGA PGSGARCHPPSTCSPSWASPG*GAKASPALPR SHGVTLLCKAQAHLCRGEDSKDASGSTSQA WEPG*GAWGMPRCQGPALGSCFCPPGTTVQ RPAKQRDKRNRHLGR
508	1858	A	3944	120	412	WCPAGTLDFPGPQEMVLLEIEVMNQLNHRNL IQLYAAIETPHEIVLFME\YECPK*W*GLGGGT TRHGASRGGVCAHSIEGGELFERIVDEDYHLT EV
509	1859	A	3949	31	392	LTKTPSPREKGRGVLSVLLMMI*KCRVIFVKIP MVFFLQNFC/RIILNVA\WTGD*PNTL*KEQRG ITFSDSKS*YKATKIKTMWYCHKNRYID/ERN RIEIPEINPCICDKIIFRKLSMTTQ
510	1860	A	3954	1013	885	FSETRACCPRLEHSGRIEAHCSLNIPGSSDPPT SASSVAATTG
511	1861	A	3956	1	1054	PPAWAPRSPLIWAPTSGRIPCRAALPWSTSSV RWQPSEKQPPPPAHRGPADSLSTAAGAAELS AEGAGKSRGSGEQDWVNRPKTVRDTLLALH QHGHSGPFESKFKKEPALTAVARTARKRKPS PEPEGEVGPPK\TTERPSRGCPHPQRGSRSP*L LHPLLCLRHHPLPHLIPTGPHRLKRPRM\P\SP MAALILVADNAGGSHASKDANQVHSTTRRN SNSPPSPSSMNQRRLGPREVGGQGAGNTGGL EPVHPASLPDSSLATSAPLCCTLCHERLEDTH FVQCPSVPSHKFCFPCSRQSIKQQGASGEVYC PSGEKCPLVGSNVPWAFMQGEIATILAGDVK VKKERDS
512	1862	A	3957	1086	3	QDRARLDCSSATSAHCNLRLPGS*DSPASASR VAGTTDTHHHTWLILGSSVQTGFDHVGQAG LELLTSGDPPISASESAGIMGMSHCVWP*SWG LSHHMAPPQGDGGRARGTPGPEQSFWNLSC H*PRCQVPS*LMTQL/FWGRHQYNPTMKRGK LRHREACSLPLPGEGEPGLQPSS\*SQNPCSSPL FHHGL*AWLWCPELLLQGQARRH*RSPPS/FK CPATLSLTAWSQTKRLRSQFLLLPWL*RAL*H PP\CHWPSRRSLGDPLLPRSQG*RDGT*ASTFC SYF*DTESHLVAQAGVQWRDLGSLQPPCPRL K\RFSRLSPPSSYTHRYVPSIILAESCISSRDRIP PSRPDRSRNSNSLSR
513	1863	A .	3961	3038	476	VALTTSMCCNKQVIVIDKIKSASIADRCGALH VGDHILSIDGTSMEYCTLAEATQFLANTTDQ VKLEILPHHQTRLALKGPDHVKIQRSDRQLT WDSWASNHSSLHTNHHYNTYHPDHCRVPAL TFPKAPPPNSPPALVSSSFSPTSMSAYSLSSLN MGTLPRSLYSTSPRGTMMRRRLKKKDFKSSL

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO; in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion  SLASSTVGLAGQVVHTETTEVVLTADPVTGF GIQLQGSVFATETLSSPPLISYIEADSPAERCG VLQIGDRVMAINGIPTEDSTFEEASQLLRDSSI TSKVTLEIEFDVAESVIPSSGTFHVKLPKKHN VELGITISSPSSRKPGDPLVISDIKKGSVAHRT GTLELGDKLLAIDNIRLDNCSMEDAVQILQQC EDLVKLKIRKDEDNSDEQESSGAIIYTVELKR YGGPLG\ITISGTEEP\FDL*IISSLTKGGLAERT GAIHIGDRIL\AINSSSLKGKPLSEAIHLLQMAG ETVTLKIKKQTDAQSASSPKKFPISSHLSDLGD VEEDSSPAQKPGKLSDMYPSHGCPSVDSAVD SWDGSA\IDTS\YGTEGT\SFQASGY\NFNTYD WRSPKQRGS\LSPVT\KPRSQTYPDVGLSYED WDRSTASGFAGAA\DSAETEQEENFWSQALE DLETCGQSGILRELEATIMSGSTMSLNHEAPT PRSPAGSDRPSFQERSSSRPHYSQTTRSNTLPS DVGRKSVTLRKMKQEIKEIMSPTPVELHKVT LYKDSDMEDFGFSVADGLLEKGVYVKNIRPA
514	1864	A	3967	833	800	GPGDLGGLKPYDRLLQVNHVRTRDFDCCLV VPLIAESGNKLDLVISRNPLASQKSIDQQSLPG D*SEQNSAFFQQPSHGGNLETREPTNTL LEKQGVSGMATKRLARQLGLIRRKSIAPANG NLGRSKSKQLFDYLIVIDFESTCWNDGKHHH SQEIIEFPAVLLNTSTGQIDSEFQAYVQPQEHPI LSEFCMELTGIKQAQVDEGVPLKICLSQFCK WIHKIQQQKNIIFATGISEPS/DF*SKIMCICYL
515	1865	A	3969	492	182	VR*RISYTY*SKHKSKGC CRFWGISTHCDTCDPLSPQTTEG**EGDLWSL DLLGPEFLARKPLFKTKTYQSTF*SISKNE/FTC PNFIIEEGTDLIF\*QVKHNPCHRLTPEEGTVQL
516	1866	A	3977		1357	NRADS KMLC/QKESNYIRLKRAKMDKSMFVKIKTLGI GAFGEVCLARKVDTKALYATKTLRKKDVLL RNQVAHVKAERDILAEADNEWVVRLYYSFQ DKDNLYFVMDYIPGGDMMSLLIRMGIFPESL ARFYIAELTCAVESVHKMGFIHRDIKPDNILID RDGHIKLTDFGLCTGFRWTHDSKYYQSGDHP RQDSMDFSNEWGDPSSCRCGDRLKPLERRAA RQHQRCLAHSLVGTPNYIAPEVLLRTGYTQL CDWWSVGVILFEMLVGQPPFLAQTPLETQM KVINWQTSLHIPPQAKLSPEASDLIIKLCRGPE DRLGKNGADEIKAHPIF*NQFDFSQ*PEDSRS AFKQFP*NHTTPTDTSNFDPVDPDKLWSDDN EEENVNDTLNGWYKNGKHPEHAFYEFTFRRF FDDNGYPYNYPKPIEYEYINSQGSEQQSDEDD QNTGSEIKNRDLVYV
517	1867	Α	3980	1358	1022	FFFKKFTQSLGFLLFSFSFLFSCFFFFHFVLFCY VFLDRVPLCHPGWSAVVQSQVT/VNLPPSWD *RCRPPH/LANLCNFCRD\SFTTLPRLVLNTWA QAIFQPQPPKVLGLQV
518	1868	A	3986	974	666	SPEMESHPITQAGVQWHFLSSLQPLPPGFK*F SCFSLPE*LGYRHVPPCLANSVFSVEMGFLH VGQAGLELLTSGDLPALASQSAGITG\SHRAR PENGFENIF
519	1869	A	3994	751	126	NQGLRHVGLCRTCLVNQMFASSILGKSHHHS LISINQGHNALWKAAG\PLPLKAGYC\QSFSPC DSLKYG\SWDEKDLTVPQRDTHKRSVLRWIS QRGK\LAVEMEEGHCLL\LPLGTECLGIK\PIV HLFSSEMGE\NRPMVG\ARHVYSNAALLSFTP

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						LRCLGGEKHKSGLHARPVIVPSLELHYDMDSI AHV\FADLLLIITLPSYYIPFC
520	1870	A	3999	882	698	QSFRLSLLSSWDYRHM*PRLANF*T\FFCRDR/ SLALLPRLVSNSWPQAILPPRPPKVLGLQT
521	1871	A	4011	1346	1178	FFF*ETVSCSAS*AGVRSHDNSSLQPPSPG\SSN PPTSASHVAGATGTHHHAWLLSV
522	1872	A	4015	2	377	QGIALLTRMGESVKHVTGGYKLRTRPLEFAA IGDYLDTFALKLGTIDRIAQRIIKEEIEYLVELR EYGPVYSTWSALEGELAEPLEGVSACIGNCST AL*ELTDDMTEDFLFVLREYILYSDSMK
523	1873	A	4018	341	19	ERVIHNQIQQAQRSPHIFNARRSS/PRPNIVELP KVKEVCKTSKS/GQVIYKGVSIRLRANFLAEP L*NRREWDEAIKVLKEKQ\FLSKMVYPANLSF GNEGDITSFPAK
524	1874	A	4020	1067	743	FFLRWSL/DSVAQAGVKWCNLGSLQAPPPGF TPFSCLSLPSSWDYRHPPPRLAN*LTNFLCF** RQGFTVLARMVLIS*PHDLPASASQSAGITGL SHCSWPTSSILS
525	1875	A	4021	781	351	QFRVIFFFLRRSHSVAQAGMQWHDHSLLQPL PPRLKQ/F/SHLSPPSIWDYRRVPPCLVNFSIFF VETGSCQPCLQLLGSSNPPASASQSAGIAGISH QGQPE*SFDIRFACVIAALRETFQCLCSASRVN NKIINRPTHPVESSF
526	1876	Α	4024	80	341	TPSSTSRGTEEQQSSKMAWQRREEKEHLNVR RSSAEDGWKADKP/VDG*TPGEDHLPTPSPFQ LHIHSSESQLHHSVKSPPSLSFRLM
527	1877	Α	4026	593	230	DFYLYPERKKRGQMMTAVSLTTRPQESVAFE DVAVYFTTKEWAIMG\PAERALYRDVMLEN YGGCGPL*CHPTSKPALVFS\LEQGKESCFSPA TGSSLSRNDWRAGWIGYLELRRYTYLS
528	1878	Α	4028	1160	242	GTSELLCIQRWNWGPAFPPRPGLALAPTLQLL VEMGSAKSVPVTPARPPPHNKHLARVADPRS PSAGILRTPIQVESSPQPGLPAGEQLEGLKHAQ DSDPRSPTLGIARTPMKTSSGDPPSPLVKQLSE VFETEDSKSNLPPEPVLPPEAPLSSELDLPLGT QLSVEEQMPPWNQTEFPSKQVFSKEEARQPT ETPVASQSSDKPSRDPETPRSS\GSMRNRWKP\ NSSKVL\GKSPLHPSCQDDNSPGTLTLRQGKA AFKPLSENVSELK\EGA\ILGTGR\LLKTEGRA WEQGQD\HDKENQHFPLVES
529	1879	Α	4039	2	366	KDMVLIMEMQSMITMKCPQYL*E*RKIPDITK CW*GCGSTGILIFC/WS*PL*KTI*QPR*FKQI*T ILTIIYSIM*EHTFHNAGV*LSDIYPRFMKGYV HTEICT*MFIAVLFVVVKTWKQF
530	1880	A	4057	358	3	LLEVNGNTIVTVFTKAQNKKNKGSRSILFKQL RKYGSRINLLKSKHDKNICTENYKT*MKEIEA /DTDKWKDILCSWIRRIHMKDILCSWIGRTHV VKISILPKVNYRFYLISIKIIMAI
531	1881	Α	4061	50	278	TQGTEEIYKISSCEWVQASFSTPLITLHDFKIY HKATVIKMVWYWHRQ*KFSKN/RIESSEIEPH IYDQFIFDKGEKIIQEKGNSFFNN/MCWKNWIF T*KR
532	1882	A	4069	19	368	NDLLENFKFWE*FKE*LENINGTVTEKETGGV YKELSSPKYSGTRQFYGQTISNFPGKIISMVY KLFQNTE/TEGRHPISLYEFRITLITIPNKDNIYL QIWMPVSLMNIVTLKCPT
533	1883	Α	4076	1	355	PIRKFTKVAG*KSNTPK*LAFLHINNEQFENKI/ ITNI/PFIIASKRIKYSGISLTKEMKDLYTETLLR KIKEDTNKWKDI/SCFWVGR/LNIVKMPK/VIC

			T 000-	T	1 = 41	
SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence	Ì	09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence			914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
1				amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
			Į	residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
-		1	1	peptide		/=possible nucleotide deletion, \=possible
			L	sequence		nucleotide insertion
						IFNAIPIKMPMMCMAKIEKNSS
534	1884	A	4088	3	1931	IIDSSTRRMESERSPLYRQLIDLGYLSSSHWNC
						GAPGQDTKAQSMLVEQSEKLRHLSTFSHQVL
ì		ľ	ì		ĺ	QTRLVDAAKALNLVHCHCLDIFINQAFDMQR
						DLQITPKRLEYTRKKENELYESLMNIANRKQE
			1			EMKDMIVETLNTMKEELLDDATNMEFKDVI
			Ι.			VPENGEPVGTREIKCCIRQIQELIISRLNQAVA
İ						NKLISSYDYLRESFYGTLERCLQSLEKSQDYS
ĺ			[		[	VHITSNYLKQILNAAYHVEVTFHSGSSVTRM
						LWEQIKQIIQRITWVSPPAITLEWKRKVAQEAI
		·				ESLSASKLAKSICSQFRTRLNSSHEAFAASLRQ
						LEAGHSGRLEKTEDLWLRVRKDHAPRLARLS
ļ		ļ	1		ļ	LESRSLODVLLHRKPKLGQELGRGQYGVVYL
						CDNWGGHFPCALKSVVPPDEKHWNDLALEF
						HYMRSLPKHERLVDLHGSVIDYNYGGGSSIA
						VLLIMERLHRDLYTGLKAGLTLETRLQIALDV
ĺ		[				VEGIRFLHSQGLVHRDIKLKNVLLDKQNRAKI
1						TDLGFCKPEAMMSGSIVGTPIHMAPELFTGK
ŀ						YDNSVDVYAFGILFWYICSGSVKLPEAFERCA
						SKDHLWNNVRRGARPERLPVFDEECWOLME
ļ		}			j	ACWDGDPLKRPLLGIVQPMLQGIMNRLCKS\
		ĺ				NSEQPNRGLDDST
535	1885	A	4090	2	417	ALMPHEANYEEIFLKTDKDMDGFESGLEVRE
333	1003	A	4090	4	417	IFLKTR/GLPSTLLAHIWALCDSKDCGKLSKD
		}			1	HFALAFHLIT\QKLIKGIDPPLVLTPEKISPSNR
i						ASLQKVTELTRKPVCIIFKGTILWRITDSIWMK
526	1006		4100	5.00	900	HNRKRIWLRA
536	1886	Α	4102	569	829	DHQK*KNIPCSWIGRINIVKMSILPKAIYRFSAI
		1				PIKIPMTFFTEI*S*NVYRTTKTQE*AKAILSKK
	1005	<u> </u>	<b> </b>			EQNLEESHYLDFK*YYRAV
537	1887	Α	4104	54	281	SIDCEHLIRRMLVLDPSKRLTIAQIKEHKWML
						IEVPVQRPVLYPQEQENEPSIGEFNEQVLRLM
						HSLGIDQQKTIE
538	1888	Α	4109	141	314	IRHIPLKIRSVVSHLKCFYKFILTFFFAGCSQPL
						VPRENITAWMNAIGLIITALPVS
539	1889	A	4111	268	1	ASRPWGHSYP*FNQQEVDTLKRPIASSEI*MM
						I*KFAT\KKSPGPYRFTAEFSHTFKEDLVPILW
		<u></u> _	<u></u>			PLFPKIYREGTLPHSFYEASITL
540	1890	A	4142	198	2064	PEPGAGRAATPWGPLFWRGRGSGRCEKAAE
						AALGDFLGLHRRTQQPAVDRLLSDASAQWR
						VRGHGGVRESGRAPQQPGRRRGRRPRKRPR
						GRWRREGCGAGGRGVCVAAWSQRSIAGNN
- 1			1			DYRLFHKMSNSHPLRPFTAVGEIDHVHILSEH
						IGALLIGEEYGDVTFVVEKKRFPAHRVILAAR
			1			CQYFRALLYGGMRESQPEAEIPLQDTTAEAFT
				1		MLLKYIYTGRATLTDEKEEVLLDFLSLAHKY
						GFPELEDSTSEYLCTILNIQNVCMTFDVASLY
						SLPKLTCMCCMFMDRNAQEVLSSEGFLSLSK
						TALLNIVLRDSFAAPEKDIFLALLNWCKHNSK
						ENHAEIMOAVRLPLMSLTELLNVVRPSGLLSP
ĺ						DAILDAIKVRSESRDMDLNYRGMLIPEENIAT
						MKYGAQVVKGELKSALLDGDTQNYDLDHG
i						
1	1	[	1			FSRHPIDDDCRSGIEIKLGQPSIINHVRILLWDR
	l			ł	I	DSRSYSYFIEVSMDELDWVRVIDHSQYLCRS
		ļ	J	Į.	}	TELEVISION AND A CONTROL OF THE PROPERTY OF TH
						WQKLYFPARVCRYIRIVGTHNTVNKIFHIVAF
						ECMFTNKTFTLEKGLIVPMENVATIADCASVI
						ECMFTNKTFTLEKGLIVPMENVATIADCASVI EGVSRSRNALLNGDTKNYDWDSGYTCHQLG
541	1891	Α	4146	282	778	ECMFTNKTFTLEKGLIVPMENVATIADCASVI

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		<u>.</u>				HAESENFAFWQDMKWKNKFWGKSLEIVPVG TVNVSLPRFGDHFEWNKVTSCIHNVLSGQRW IEHYGEVLIRNTQDSSCHCKITFCKAKYWSSN VHEVQGAVLSRSGRVLHRLFGKWHEGLYRG PTPGGQCIWKP
542	1892	A	4147	44	433	SVDAYVCNDIVFSYRTTITLLEGA*LTHRYVA QDPKQGQLRSLHLTCDSAPAGSQGTWSTSCR INHLIFRGGAQITFLATFDDSPKAVLGDRLLLT ANVSSENNTPRTSKTTFQLELSVKDAVYTVV SSH
543	1893	A	4153	678	11	TISYPQCLTQMYFLISFANVDTFLLPIMALDH YVAICSALQ*CSIITP/ELCQGLPVLA*AGSSLIS PVHTVIMSRLAFCSSAQISHFYRDAYLLMKIA CSHT*\NQHVFLGAVVLFLAPCALILVSYIRIA AAILRIPSPTRRKACSICSSHLSLVTLFYGTV LGICI*PPDSFSAQDAIATIMYTVVTSMLNPFIY SLMNKEVQEAVRRLFSRGSHSSWCW
544	1894	A	4158		538	LLYAQAGVQ*LNLSSLQPQPAGLKQSSHPSLP SSWDYRYSTPHPANFFVEMEFHHVAQAGLEL LGSGDLPTSTSHSAGITGV\SHHAPPRLISSEGS LLGHLLCLPMVFPLLCVFVLISSSLAGEEAAG LRVQKLWPAVVLSHLPVCWFHCSGIWSEVIE LKVGREGHVLPWQAHVVEF
545	1895	A	4160	1	412	HPLGLGLVPSEIFSPQDKKAADGSILAPARGE DLEAGLKGSFMDGRLQASVSVFRIQRVGSAM QDTASAMPCLPYYPTSHCFMAGGKSRSQGW ELELSGEPAPGWQVLAGYTYTQARYLRDASE ANVGQPLRPVDPR
546	1896	A	4174	1252	1190	FFQVFIFLFIFFKTEFHSCCPGAVQWHDLDSL QPPPPRFKGFSCLSLPSSWDYRHAPAHPANFV FLVETGFLHV\GQ\ASLELPTSGDTPAS\ASQSA GITGVSHHA*PRASGRRCW
547	1897	A	4176	3029		AGPDGLAAPASCQGARGQTRVPGAFSWLAP GSHHASEGLAPGVPPAGGVSAQELTAPPQEG WGLGAPPAAPRPESDEKRAGSDAVRSFSRGA RDSLGQRRLGGTRGAGPAGKGAQRTMGPAS GFHSFPRPHQEPSPRSSCWQHLLWHCPWPQ PSRLPRLTPAQLLQGPGVLAAPPGP*HVPGFL AQSPWPLPSGPRSP*DPLHQGALVPLPQGGSP HTAPHCLPSVLSPAIQQPLLPTAST/SSRSPPAS TMAPIPSALAVWEPAGSSPQLSSAPADSS/PLP ALPKVLPPWTQKPLLGCLCQSPLPLLSPPDQI/ RCPPACSPAAASSFSFESQPCPSAPSKASPAPA ALIVGPHHPP*SQQPQSQSVHPHGPGGPQPPL AASSLFWMFCQPPPPHPQFLWHRPLPVTGKA LAS/PLCFRPAPGSLRQTPLPPQFHIPRPGLSAP/ PPPASGTSDSSDSRSPSASAARVWPPA\SPPPP AARHRPHPPEYFLSPCPFSCGFPRLLGRPRRPQ ALQTPRAWDLPPGSSPAPLCSGPELP*APPPLP PFPRVA*LGSGHPPSAQVPGLW*RCV*GHPIP RPVGHS*SGPPHSPPL*APPQAWPLELPPSRQC LQPLHLRAAQPLDPCCSLSPPGPPLPVPALPS WPGRP*SPSPASSQPPYHAGLPGPQSSPLPPGL PQLPSLRSGSQQPLLFFQCPGPGAVWGKGSPQ PLSPHPPPP/ARTQTFPVASRSLSPGTAPYSVCL TPSRSASSLPEVVLASSLPKIPQSSGS\PLGPTSP MP*CFHRPSPPLP/LSSPFPA\LRPQAPQFPLHLP P*PAPSPGCPLPPLAQQHQPSPPSPHARSTLT PPLWPSLALLP*PLPPPPPVPSFSASLLCSLPAH

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion GTPASPGLGRSCLGKPQTLPWISFWPPSGRLA PGTWQPW/PVSPAPLSCLSAWDPWELPSPQPQ VCSTAELPTSCLLSSPGP\PAFOPPRFGCL*GPP
548	1898	A	4180	2369	844	GPPGLPPLQSSLSFPPPPPPPVPQPPAPPALQWG LHLPGGRTK RIHREEDFQFILKGIARLLSNPLLQTYLPNSTK
	1090			2309	0.74	KIQFHQELLVLFWKLCDFNKVGQPRGALQGD GEQLPQ*PGGRDSVRLRGVGQSCPSLELSPLG PSPHP*KFLFFVLKSSDVLDILVPILFFLNDAR ADQSRVGLMHIGVFILLLLSGECNFGVRLNKP YSIRVPMDIPVFTGTHADLLIV\VFHKIITSGHQ RLQPLFDCLLTIVVNVSPYLKSLSMVTANKLL HLLEAFSTTWFLFSAAQNHHLVFFLLEVFNNI IQYQFDGNSNLVYAIIRKRSIFHQLANLPTDPP TIHKALQRRRTPEPLSRTGSQGGAPPWRAPA PLPLQSQAPSRPVWWLLQALTS*PRSPRCQR MAPCGPWNLSPSRAWRMAARLRGSPARHGG SSGDRP/HSSASGQWSPTPEWVLSWKSKLPLQ TIMRLLQVLVPQVEKICIDKGLTDESEILRFLQ HGTLVGLLPVPHPILIRKYQANSGTAMWFRT YMWGVIYLRNVDPPVWYDTDVKLFEIQRV
549	1899	A	4191	858	321	LPWQRLGVLLSRGKMAVTGWLESLRTAQKT ALLQDGRRKVHYLFPDGKEMAEEYDEKTSE LLVRKWRVKSALGAMGQWQLEVGDPAPLG AGNLGPELIKESNANPIFMRKDTKMSFQWRIR NLPYPKDVYSVSVDQKERCIIVRTTNKKYYK KFSIPDLDRHQLPLDDALLSFA\TPTAP
550	1900	A	4192	1	1980	IRHTGSDIAGVCGWLLLSGPCGVGLDLDSRLL GASAMRRSEVLAEESIVCLQKALNHLREIWE LIGIPEDQRLQRTEVVKKHIKELLDMMIAEEE SLKERLIKSISVCQKELNTLCSELHVEPFQEEG ETTILQLEKDLRTQVELMRKQKKERKQE\LKL LQEQDQELC\EILCMPHYDIDSASVPSLEELNQ FRQHVTTLRETKASRREEF/VSSIKRQIILCME ELDHTPDTSFERDVVCEDEDAFCLSLENIAT\L QKLLRQ\LEMQKSQNEAVCEG\LRTQI\RELW DRLQIPEEEREAVATIMSGSKAKVRK\ALQ\LE VDRLEELEKCKTMKKVIEARVELVQYWDQC FYSQEQRQAFAPFCAEDYTESLLQLHDAEIVR LKNYYEVHKELFEGVQKWEETWRLFLEFER KASDPNFTNRGGNLLKEEKQRAKLQKMLP KLEELKARIELWEQEHSKAFMVNGQKFME YVAEQWEMHRLEKERAKQERQLKNKKQTET EMLYGSAPRTPSKRRGLAPNTPGKARKLNTT TMSNATANSSIRPIFGGTVYHSPVSRLPPSGSK PVAASTCSGKKTPRTGRHGANKENLELNGSI LSGGYPGSAPLQRNFSINSVASTYSEFADPSLS DSSTVGLQRELSKASKSVGRAVITATIV
551		A	4194	3	1008	AWHEGLVSSPAIGAYLSASYGDSLVVLVATV VALLDICFILVAVPESLPEKMRPVSWGAQISW KQADPFASLKKVGKDSTVLL\ICITVCLSYLPE AG\QYSSFF\LYLR\QVIGFG\TVKIAAFIAMVGI LSIVAQTAFLSILMRSLGNKNTVLLGLGFQML QLAWYGFGSQAWMMWAAGTVAAMSSITFP AISALVSRNAESDQQGVAQGIITGIRGLCNGL GPALYGFIFYMFHVELTELGPKLNSNNVPLQ GAVIPGPPFLFGACIVLMSFLVALFIPEYSKAS GVQKHSNSSSGSLTNTPERGSDEDIEPLLQDS SIWELSSFEEPGNQCTEI.

SEO ID	SEO ID	Mot	Lego	Destina	D. 1:4-41	· (A. Al. i C. G
SEQ ID NO: of	SEQ ID NO: of	Met hod	SEQ ID NO:	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide	nou	in in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence		ĺ	914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
		]		amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
				peptide	•	/=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion
552	1902	A	4197	2	14302	ARPPPAPGSRQQKQKAAPGAAAAAELRGAR
						EPAPARRRGTMADGGEGEDEIQFLRTDDEVV
}						LQCTATIHKEQQKLCLAAEGFGNRLCFLESTS
						NSKNVPPDLSICTFVLEQSLSVRALQEMLANT
						VEKSEGQVDVEKWKFMMKTAQGGGHRTLL
			}			YGHAILLRHSYSGMYLCCLSTSRSSTDKLAFD
						VGLQEDTTGEACWWTIHPASKQRSEGEKVR
						VGDDLILVSVSSERYLHLSYGNGSLHVDAAF QQTLWSVAPISSGSEAAQGYLIGGDVLRLLH
						GHMDECLTVPSGEHGEEQRRTVHYEGGAVS
						VHARSLWRLETLRVAWSGSHIRWGQPFRLR
						HVTTGKYLSLMEDKNLLLMDKEKADVKSTA
						FTFRSSKEKLDVGVRKEVDGMGTSEIKYGDS
						VCYIQHVDTGLWLTYQSVDVKSVRMGSIQR
			[			KAIMHHEGHMDDGISLSRSQHEESRTARVIRS
						TVFLFNRFIRGLDALSKKAKASTVDLPIESVSL
						SLQDLIGYFHPPDEHLEHEDKQNRLRALKNR
						QNLFQEEGMINLVLECIDRLHVYSSAAHFAD
			}			VAGREAGESWKSILNSLYELLAALIRGNRKN
				İ		CAQFSGSLDWLISRLERLEASSGILEVLHCVL
						VESPEALNIIKEGHIKSIISLLDKHGRNHKVLD
}						VLCSLCVCHGVAVRSNQHLICDNLLPGRDLL
						LQTRLVNHVSSMRPNIFLGVSEGSAQYKKWY YELMVDHTEPFVTAEATHLRVGWASTEGYSP
-						YPGGGEEWGGNGVGDDLFSYGFDGLHLWSG
1						CIARTVSSPNQHLLRTDDVISCCLDLSAPSISF
						RINGQPVQGMFENFNIDGLFFPVVSFSAGIKV
						RFLLGGRHGEFKFLPPPGYAPCYEAVLPKEKL
						KVEHSREYKQERTYTRDLLGPTVSLTQAAFT
						PIPVDTSQIVLPPHLERIREKLAENIHELWVMN
İ						KIELGWQYGPVRDDNKRQHPCLVEFSKLPEQ
						ERNYNLQMSLETLKTLLALGCHVGISDEHAE
J	j				j	DKVKKMKLPKNYQLTSGYKPAPMDLSFIKLT
						PSQEAMVDKLAENAHNVWARDRIRQGWTY
]						GIQQDVKNRRNPRLVPYTPLDDRTKKSNKDS
ļ					ļ	LREAVRTLLGYGYNLEAPDQDHAARAEVCS
						GTGERFRIFRAEKTYAVKAGRWYFEFETVTA
						GDMRVGWSRPGCQPDQELGSDERAFAFDGF KAQRWHQGNEHYGRSWQAGDVVGCMVDM
						NEHTMMFTLNGEILLDDSGSELAFKDFDVGD
						GFIPVCSLGVAQVGRMNFGKDVSTLKYFTIC
1					ĺ	GLQEGYEPFAVNTNRDITMWLSKRLPQFLQV
						PSNHEHIEVTRIDGTIDSSPCLKVTQKSFGSQN
						SNTDIMFYRLSMPIECAEVFSKTVAGGLPGAG
	ľ	ĺ			1	LFGPKNDLEDYDADSDFEVLMKTAHGHLVP
		ļ				DRVDKDKEATKPEFNNHKDYAQEKPSRLKQ
						RFLLRRTKPDYSTSHSARLTEDVLADDRDDY
1	ľ		ľ		1	DFLMQTSTYYYSVRIFPGQEPANVWVGWITS
					1	DFHQYDTGFDLDRVRTVTVTLGDEKGKVHE
					İ	SIKRSNCYMVCAGESMSPGQGRNNNGLEIGC
1	1	ľ	1		ľ	VVDAASGLLTFIANGKELSTYYQVEPSTKLFP
ļ		ļ	j	1	ļ	AVFAQATSPNVFQFELGRIKNVMPLSAGLFKS
	}		j			EHKNPVPQCPPRLHVQFLSHVLWSRMPNQFL
-	İ					KVDVSRISERQGWLVQCLDPLQFMSLHIPEEN
1				J		RSVDILELTEQEELLKFHYHTLRLYSAVCALG NHRVAHALCSHVDEPQLLYAIENKYMPGLLR
	1					AGYYDLLIDIHLSSYATARLMMNNEYIVPMT
	-					
	-					
	-	•				EETKSITLFPDENKKHGLPGIGLSTSLRPRMQF
	-					

SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide	1	in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence			914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
dones			71.1	amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
i l				residue of	sequence	
1					sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
1	ì	1	Ì	peptide	ĺ	/=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion
						MGIFHNEDLKHILQLIEPSVFKEAATPEEESDT
	-					LEKELSVDDAKLQGAGEEEAKGGKRPKEGLL
	}					QMKLPEPVKLQMCLLLQYLCDCQVRHRIEAI
		]				VAFSDDFVAKLQDNQRFRYNEVMQALNMSA
1						ALTARKTKEFRSPPQEQINMLLNFKDDKSECP
						CPEEIRDQLLDFHEDLMTHCGIELDEDGSLDG
						NSDLTIRGRLLSLVEKVTYLKKKQAEKPVES
						DSKKSSTLQQLISETMVRWAQESVIEDPELVR
	}	1				AMFVLLHRQYDGIGGLVRALPKTYTINGVSV
						EDTINLLASLGQIRSLLSVRMGKEEEKLMIRG
						LGDIMNNKVFYQHPNLMRALGMHETVMEV
		[ .				MVNVLGGGESKEITFPKMVANCCRFLCYFCR
1	1					
		!		·		ISRQNQKAMFDHLSYLLENSSVGLASPAMRG
	ĺ	•				STPLDVAAASVMDNNELALALREPDLEKVVR
						YLAGCGLQSCQMLVSKGYPDIGWNPVEGER
						YLDFLRFAVFCNGESVEENANVVVRLLIRRPE
	1					CFGPALRGEGGNGLLAAMEEAIKIAEDPSRD
						GPSPNSGSSKTLDTEEEEDDTIHMGNAIMTFY
		1				SALIDLLGRCAPEMHLIHAGKGEAIRIRSILRS
						LIPLGDLVGVISIAFQMPTIAKDGNVVEPDMS
	ł	l				AGFCPDHKAAMVLFLDRVYGIEVQDFLLHLL
					'	EVGFLPDLRAAASLDTAALSATDMALALNRY
						LCTAVLPLLTRCAPLFAGTEHHASLIDSLLHT
						VYRLSKGCSLTKAQRDSIEVCLLSICGQLRPS
						MMQHLLRRLVFDVPLLNEHAKMPLKLLTNH
1	[	ĺ 1				YERCWKYYCLPGGWGNFGAASEEELHLSRK
						LFWGIFDALSQKKYEQELFKLALPCLSAVAG
						ALPPDYMESNYVSMMEKQSSMDSEGNFNPQ
						PVDTSNITIPEKLEYFINKYAEHSHDKWSMDK
}						LANGWIYGEIYSDSSKVQPLMKPYKLLSEKE
						KEIYRWPIKESLKTMLARTMRTERTREGDSM
1						ALYNRTRRISQTSQVSVDAAHGYSPRAIDMS
			1			NVTLSRDLHAMAEMMAENYHNIWAKKKKM
						ELESKGGGNHPLLVPYDTLTAKEKAKDREKA
			1			QDILKFLQINGYAVSRGFKDLELDTPSIEKRFA
1 1						YSFLQQLIRYVDEAHQYILEFDGGSRGKGEHF
						PYEQEIKFFAKVVLPLIDQYFKNHRLYFLSAA
						SRPLCSGGHASNKEKEMVTSLFCKLGVLVRH
]					]	RISLFGNDATSIVNCLHILGOTLDARTVMKTG
					j	
]						LESVKSALRAFLDNAAEDLEKTMENLKQGQF
						THTRNQPKGVTQIINYTTVALLPMLSSLFEHI
1 1						GQHQFGEDLILEDVQVSCYRILTSLYALGTSK
						SIYVERQRSALGECLAAFAGAFPVAFLETHLD
						KHNIYSIYNTKSSRERAALSLPTNVEDVCPNIP
		1				SLEKLMEEIVELAESGIRYTOMPHVMEVILPM
1						LCSYMSRWWEHGPENNPERAEMCCTALNSE
j l						HMNTLLGNILKIIYNNLGIDEGAWMKRLAVF
] [				[	ſ	SOPIINKVKPOLLKTHFLPLMEKLKKKAATVV
						SEEDHLKAEARGDMSEAELLILDEFTTLARDL
						YAFYPLLIRFVDYNRAKWLKEPNPEAEELFR
				ļ	l	MVAEVFIYWSKSHNFKREEQNFVVQNEINN
1						MSFLITDTKSKMSKAAVSDQERKKMKRKGD
						RYSMQTSLIVAALKRLLPIGLNICAPGDQELIA
						LAKNRFSLKDTEDEVRDIRSNIHLQGKLEDP
			1			AIRWQMALYKDLPNRTDDTSDPEKTVERVL
			- 1			
]			1			DIANVLFHLEQKSKRVGRRHYCLVEHPQRSK
1	ı	ı J	į		}	KAVWHKLLSKQRKRAVVACFRMAPLYNLPR
1						
{ I			ļ			HRAVNLFLQGYEKSWIETEEHYFEDKLIEDLA
						HRAVNLFLQGYEKSWIETEEHYFEDKLIEDLA KPGAEPPEEDEGTKRVDPLHQLILLFSRTALT
			•			

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
			<i>f</i>			EEEVKSFEEKEMEKQKLLYQQARLHDRGAA EMVLQTISASKGETGPMVAATLKLGIAILNGG NSTVQQKMLDYLKEKKDVGFFQSLAGLMQS CSVLDLNAFERQNKAEGLGMVTEEGSGEKV LQDDEFTCDLFRFLQLLCEGHNSDFQNYLRT QTGNNTTVNIIISTVDYLLRVQESISDFYWYY SGKDVIDEQGQRNFSKAIQVAKQVFNTLTEYI QGPCTGNQQSLAHSRLWDAVVGFLHVFAHM QMKLSQDSSQIELLKELMDLQKDMVVMLLS MLEGNVVNGTIGKQMVDMLVESSNNVEMIL KFFDMFLKLKDLTSSDTFKEYDPDGKGVISK RDFHKAMESHKHYTQSETEFLLSCAETDENE TLDYEEFVKRFHEPAKDIGFNVAVLLTNLSEH MPNDTRLQTFLELAESVLNYFQPFLGRIEIMG SAKRIERVYFEISESSRTQWEKPQVKESKRQFI FDVVNEGGEKEKMELFVNFCEDTIFEMQLAA QISESDLNERSANKEESEKERPEEQGPRMAFF SILTVRSALFALRYNILTLMRMLSLKSLKKQM KKVKKMTVKDMVTAFFSSYWSIFMTLLHFV ASVFRGFFRIICSLLLGGSLVEGAKKIKVAELL ANMPDPTQDEVRGDGEEGERKPLEAALPSED LTDLKELTEESDLLSDIFGLDLKREGGQYKLIP HNPNAGLSDLMSNPVPMPEVQEKFQEQKAK EEEKEEKEETKSEPEKAEGEDGEKEEKAKED KGKQKLRQLHTHRYGEPEVPESAFWKKIIAY QQKLLNYFARNFYNMRMLALFVAFAINFILL FYKVSTSSVVEGKELPTRSSSENAKVTSLDSS SHRIIAVHYVLEESSGYMEPTVRILPILHTVISF FCIIGYYCLKVPLVIFKREKEVARKLEFDGLYI TEQPSEDDIKGQWDRLVINTQSFPNNYWDKF VKRKVMDKYGEFYGRDRISELLGMDKAALD
553	1903	A	4199	31	767	FSDAREKKKPKKDSSLSAVLNSIDVKYQMW KLGVVFTDNSFLYLAWYMT LPELNGRGAGLRRAEPSERGGGAERTQQVAA
						LPLSHGHSHGGGGCRCAAER/VGAARGSAAC AYGLYLRIDKGRLQCLNESREGSGRGVFKPW ERAD\DRSKFVESDADEELLFNIPFTG\HVKLK GIIIMGEDDDSHPSEMRLYKNIPQMSFDDTER EPDQTFSLNRDLTGELEYATKISRFSNVYHLSI HISKNFGADTTKVFYIGLRGEWTELRRHEVTI CNYEASANPADHRVHQVTPQTHFIS
554	1904	A	4200	1	961	GIPCTEMGNFDNANVTGEIEFAIHYCFKTHSL EICIKACKNLAYGEEKKKKCNPYVKTYLLPD RSSQGKRKTGVQRNTVDPTFQETLKYQVAPA QLVTRQLQVSVWHLGTLARRVFLGEVIIPLAT WDFEDSTTQSFRWHPLRAKADKYEDSVPQS NGELTVRAKLVLPSRTRKLQEAQEGTDQPSL HGQLCLVVLGAKNLPVRPDGTLNSFVKGCLT LPDQQKLRLKSPVLRKQACPQWKHSFVFSGV TPAQLRQSSLELTVWDQALFGMNDRLLGGT\ RLGSKGDTAVGGDACSQSKLQWQKVLSSPN LWTDMTLVLH
555	1905	A	4211	331	2419	KENKKARNIRMNQSRSRSDGGSEETLPQDH NHHENERRWQQERLHREEAYYQFINELNDE DYRLMRDHNLLGTPGEITSEELQQRLDGVKE QLASQPDLRDGTNYRDSEVPRESSHEDSLLE WLNTFRRTGNATRSGQNGNQTWRAVSRTNP NNGEFRFSLEIHVNHENRGFEIHGEDYTDIPLS DSNRDHTANRQQRST\SPVARRTRSQTSVNFN GSSSNIPRTRLASRGQNPAEGSFSTLGRLRNGI

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  GGAAGIPRANASRTNFSHTNQSGGSELRQRE GQRFGAAHVWENGARSNVTVRNTNQRLEPI RLRSTSNSRSRSPIQRQSGTVYHNSQRESRPV QQTTRRSVRRGRTRVFLEQDRERERGTAY TPFSNSRLVSRITVEEGEESSRSSTAVRHPTIT LDLQVR\RIRPGENRDRDSIANRTRSRVGLAE NTVTIESNSGGFRRTISRLERSGIRTYVSTITVP LRRISENELVEPSSVALRSILRQIMTGFGELSSL MEADSESELQRNGQHLPDMHSELSNLGTDN NRSQHREGSSQDRQAQGDSTEMHGENETTQP HTRNSDSRGGRQLRNPNNLVETGTLPILRLAH FFLLNESDDDDRIRGLTKEQIDNLSTRHYEHN SIDSELGKICSVCISDYVTGNKLRQLPCMHEF HIHCIDRWLSENCTCPICRQPVLGSNIANNG
556	1906	Α	4212	3	462	LQRQRQHPAAAPAVPVRCFTFCFTDIVIMPKR KSPENTEGKDGSKVTKQEPTRRSARLSAKPA PPKPEPKPRKTSAKKEPGAKISRGAKGKKEEK QEAGKEGTAPSENGETKAEEIHISRSTVNVST SRGTPPSTLSVKGQIETVRVKGTEN
557	1907	A	4213	774	507	ARRFSCLTLQTSWGHRH\GPPRP\ANFVFLVET GFLHIGQAGHKLPTSGDPPASASQSARITGMS HRTWFLASFLIDSCKNFIVYKIMYTL
558		A	4225	3	1253	TYRHAEREHPETSSATKVSYDYRHKRPKLLD GDQDFSDGRTQKYCKEEDRKYSFQKGPLNRE LDCFNTGRGRETQDGQVKEPFKPSKKDSIAC TYSNKNDVDLRSSNDKWKEKKKKEGDCRKE SNSSSNQLDKSQKLPDVKPSPINLRKKSLTVK VDVKKTVDTFRVASSYSTERQMSHDLVAVG RKSENFHPVFEHLDSTQNTENKPTGEFAQEIIT IIHQVKANYFPSPGITLHERFS\KMADIHKADV NEIPLNSDPEIHRRIDMSLAELQSKQAVIYESE QTLIKIIDPNDLRHDIERRKERLQNEDEHIFHI ASAAERDDQNSSFSKNYTTQRKDIITHKPFEV EGNHRNTRVRPFKSNFRGGRCQPNYKSGLVQ KSLYIQAKYQRLRFTGPRGFITHKFRERLMRK KKVP
559	1909	A	4235	1	323	KFSIPFFLRWSFTLV\PRLEGNDMISVHCNLGL LGLSHSPASASQVGGITGTQHHTGLIFGFLIET EFHHVGQAGLELLTSGDPPALAFQSAGITGVS HHAWLQVLNS
560	1910	A	4246	2	1569	TLSLLERVLMKDIVTPVPQEEVKTVIRKCLEQ AALVNYSRLSEYAKIEGKKREMYELPVFCLA SQVMDLTIQNQKDAENVGRLITPAKKLEDTIR LAELVIEVLQQNEEHHAEAFAWWSDLMVEH AETFLSLFAVDMDAALEVQPPDTWDSFPLFQ LL\NDFLRTGLLICGNGK\FHKHLQDLFAPLVV R/YMWDLDGSSPIAQSIHRGLLSRESWEPVNN GSGTSEDLFWKLDALQTFIRDLHWPEEEFGK HLEQRLKLMASDMIESCVKRTR\IAFEVKLQK TSSIQQIFRVPQFNMAPCFNVMGLMAKGSIQP KL\CSMEMGQEFAKMWHQYHSKIDELIEETV KEMITLLVAKFVTILEGVLAKLSRYDEGTLFS SFLSFTVKAASKYVDVPKPGMDVADAYVTF VRHSQDVLRDKVNEEMYIERLFDQWYNSSM NVICTWLTDRMDLQLHIYQLKTLIRMVKKTY RDFRLQGVLDSTLNSKTYETIRNRLTVEEATA SVSEGGGLQGISMKDSDEEDEEDD
561	1911	A	4257	1300	654	SELVQFLLIKDQKKIPIKRADILKHVIGDYKDI FPDLFKRAAERLQYVFGYKLVELEPKSNTYIL

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion INTLEPVEEDAEMRGDQGTPTTGLLMIVLGLI FMKGNTIKETEAWDFLLAL\GVYPTKKHLIFG
						DPKKLITEDFVRQRYLEYRRIPHTDPVDYEFQ WGPRTNLETSKMKVLKFVAKVHNQDPKDW PAQYCEALADEENRARPQPSGPAPSS
562	1912	A	4260	1	1498	MVTWLYRFLPTSNMAAKLRSLLPPDLRLQF WLHARLQKCFLSRGCGSYCAGAKASPLPGK MAMGLMCGRRELLRLLQSGRRVHSVAGPSQ WLGKPLTTRLLFPAAPCCCRPHYLFLAASGPR SLSTSAISFAEVQVQAPPVVAATPSPTAVPEV ASGETĄDVVQTAAEQSFAELGLGSYTPVGLI QNLLEFMHVDLGLPWWGAIAACTVFARCLIF PLIVTGQREAARIHNHLPEIQKFSSRIREAKLA GDHIEYYKASSEMALYQKKHGIKLYKPLILPV TQAPIFISFFIALREMANLPVPSLQTGGLWWF QDLTVSDPIYILPLAVTATMWAVLELGAETG VQSSDLQWMRNVIRMMPLITLPITMHFPTAV FMYWLSSNLFSLVQVSCLRIPAVRTVLKIPQR VVHDLDKLPPREGFLESFKKGWKNAEMTRQ LREREQRMRNQLELAARGPLRQTFTHNPLLQ PGKDNPPNIPSS\SSSSSSKPKSKYPWHDTLG
563	1913	A	4265	623	116	MGGLAPTQTLEPT\REYQNTQLSVSYLLPEQN THGTRRTLSSGPSNNLPLPLSSSATMPSMQCK HRSPNGGLFRQSPVK/TPPIPMSFQPVPGGV\L PRGSGNPPHGTSILTAPPALLPHPPTHPTQQSF LIQENNNTNHTHSHTHTYTETLSFFLYICVNN DRMEWGKSVF
564	1914	A	4270	3	368	ILKRKLSSLNSEVSTIQNTRMLAFKATAQLFIL GCTWCLGLLQVGPAAQVMAYLFTIINSLQGF FIFLVYCLLS\QQVQKQYQKWFREIVKSKSES ETYTLSSKMGPDSKPSEGDVFPRTSE
565	1915	A	4288	83	406	RNSRPLWCSPPASQPRQAPVSQSCCCPLPSSSS PPSALLAPTKPRALGTLRLYECSPELCTTMLP PAWLLMLCQAPRPQDPDPRLTQPEKSLQEAP GQTGASRTPRT
566	1916	A	4298	1041	229	LNSSQKLACLIGVEGGHSLDSSLSVLRSFYVL GVRYLTLTFTCSTPWAESSTKFRHHMYTNVS GLTSFGEKVVEELNRLGMMIDLSYASDTLIRR VLEVSQAPVIFSHSAARAVCDNLLNVPDDILQ LLKKNGGIVMVTLSMGVLQCNLLANVSTVA DHFDHIRAVIGSEFIGIGGNYDGTGRFPQGL\(\text{E}\) DVSTYPVLIEELLSRS\(\text{WSEEELQGVLRGNLLR}\) VFRQVEKVREESRAQSPVEAEFPYGQLSTSCH FHLGASEWTPRLLIWR
567	1917	A	4299	1	1106	GATPLGSVGGRTGKMDAATLTYDTLRFAEFE DFPETSEPVWILGRKYSIFTEKDEILSDVASRL WFTYRKNFPAIGGTGPTSDTGWGCMLRCGQ MIFAQALVCRHLGRDWRWTQRKRQPDSYFS VLNAFIDRKDSYYSIHQIAQMGVGEGKSIGQ WYGPNTVAQVLKKLAVFDTWSSLAVHIAMD NTVVMEEIRRLCRTSVPCAGATAFPADSDRH CNGFPAGAEVTNRPSPWRPLVLLIPLRLGLTD INEAYVETLKHCFMMPQSLGVIGGKPNSAHY FIGYVGEELIYLDPHTTQPAVEPTDGCFIPDES FHCQHPPCRMSIAELDPSIAVVRGGHLSTQAF GAECCLGMTRKTFGFLRFFFSMLG
568	1918	A	4300	2012	1843	SRKFLTITPIVLYFLTSFYTKYDQIHFVLNTVS LMSVLIPKLPQLHGVRIFGINKY
569	1919	Ā	4302	186	531	WTFCLFL/WWVPESARWLLTQGHVKEAHRY

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion LLHCARLNGRPVCEDSFSQEVRVNVCVSMHI
						CVWWGVGCVKCLPPRAHHIWQEKPLGPHRT VTESKLEAEGKTKEKAREKERKKKS
570	1920	A	4308	3		RSGQGKYYGLIGRRFQQMDVLEGLNLITIS GKRNKLRVYYLSWLRNKILHNDPEVEKKQG WTTVGDMEGCGHYRVVKYERIKFLVIALKSS VEVYAWAPKPYHKFMAFKSFADLPHRPLLV DLTVEEGQRLKVIYGSSAGFHAVDVDSGNSY DIYIPVHIQSQITPHAIIFLPNTDGMEMLLCYE DEGVYVNTYGRIIKDVVLQWGEMPTSVAYIC SNQIMGWGEKAIEIRSVETGHLDGVFMHKRA QRLKFLCERNDKVFFASVRSGGSSQVYFMTL NRNCIMNW
571	1921	A	4309	9	524	ASREMDVTKVCGEMRYQLNKTNMEKDEAE KEHREFRAKTNRDLEIKDQEIEKLRIELDESK QHLEQEQQKAALAREECLRLTELLGESEHQL HLTRQEKDSIQQSFSKEAKAQALQAQQREQE LTQKIQQMEAQHDKTENEQYLLLTSQNTFLT KLKEECCTLAKKLEQISQ
572	1922	A	4318	1	1119	GATPLGSVGGRTGKMDAATLTYDTLRFAEFE DFPETSEPVWILGRKYSIFTEKDEILSDVASRL WFTYRKNFPAIGGTGPTSDTGWGCMLRCGQ MIFAQALVCRHLGRDWRWTQRKRQPDSYFS VLNAFIDRKDSYYSIHQIAQMGVGEGKSIGQ WYGPNTVAQVLKKLAVFDTWSSLAVHIAMD NTVVMEEIRRLCRTSVPCAGATAFPADSDRH CNGFPAGAEVTNRPSPWRPLVLLIPLRLGL\T DINEAYV\ETL\KHCFHGWPQFPG/VVHREGK PNSAHYFIGYVGEELIYLDPHTTQPAVEPTDG CFIPDESFHCQHPPCRMSIAELDPSIAVVRGGH LSTQAFGAECCLGMTRKTFGFLRFFFSMLG
573	1923	A	4333	363	1066	GGVPVGLASKPFQILYGHTNEVLSVGISTELD MAVSGSRDGTVIIHTIQKGQYMRTLRPPCESS LFLTIPNLAISWEGHIVVYSSTEEKTTLKÆRM HYICFSINGKYLGSQILKEQVSDICIIGEHIVTG SIQGFLSIRDLHSLNLSINPLAMRLPIHCVCVT KEYSHILVGLEDGKLIVVGVGKPAEVKPSISN FISHAVGDYFGSPSFQLIEKSPLGINKLKAKFD FSKGSK
574	1924	A	4346	359	1234	MDTLEEVTWANGSTALPPPLAPNISVPHRCLL LLYEDIGTSRVRYWDLLLLIPNVLFLIFLLWK LPSARAKIRITSSPIFITFYILVFVVALVGIARA VVSMTVSTSNAATVADKILWEITRFFLLAIEL SVIILGLAFGHLESKSSIKRVLAITTVLSLAYSV TQGTLEILYPDAHLSAEDFNIYGHGGRQFWL VSSCFFFLVYSLVVILPKTPLKERISLPSRRSFY VYAGILALLNLLQGLGSVLLCFDIIEGLCCVD ATTFLYFSFFAPLIYVAFLRGFFGSEPKILF
575	1925	A	4360	2038	1512	GCWRHPWLASQRDCLDCRIQLAEKFVKAV SKPSRPDMNPIRVKEVYRLEEMEKIFVRLEM KIIKGSSGTPKLSYTGRDDRHFVPMGLYIVRT VNEPWTMGFSKSFKKKFFYNKKTKDSTFDLP ADSIAPFHICYYGRLFWEWGDGIRVHDSQKP QDQDKLSKEDVLSFIQMHRA
576	1926	A	4365	69	500	QVEGRQGREVKRTAWRISPVWRPARCRRRST PQP/PE/PGAQQQERHRQGEAPMQALDPRAEP GPQAQSHAACQPEPEPPRVLLDPTAARGGVQ GRP/GLSRHPGLAPHPQTHTPWPQSGRLPCAS EPLPLGGIRPTPGLEPKGRDLM

OF C	OFO TO	N #	CEO	I D 11 1	l m. it is the	
SEQ ID NO: of	SEQ ID NO: of	Met hod	SEQ ID NO:	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
nucl-	peptide	nou	in NO.	beginning nucleotide	nucleotide location	D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence	1	09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence	acrico		914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
			7	amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
		l '		residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
		1		peptide		/=possible nucleotide deletion, \=possible
			,	sequence		nucleotide insertion
577	1927	A	4366	785	502	SAPPKKKNGVLFLSPRLKSSGAIWVHSTPTLW
						ASSNSRASTPKVAGITGARPHARIIFVFLIEMG
						FHNVGQAGL/DTLTLVICPPQPPKLLGLQM
578	1928	A	4367	1	221	FFFFLKKSRCVTQAGVQG\PISLHPPPPGFKRF
						SRLSLLSSWDYRHP/HAANFCIFSRDG\VSPYW
		İ				SGWSRTPDLR
579	1929	A	4383	1	224	FETESHSVTQAGMQWHNLGSLQPMP/PGLKR
		1				FSCLRLQSSWDHRHAPPHLAHFCIFSRDGVSP
						CWPGWSSTPDLK
580	1930	Α	4397	410	94	SRLKPYSTNVTAKKLPATNIPNLDCFTAKLYQ
						\VFKKGI\IHILHELFQNKEEGAFPNS/FYEASFT
						LRPKSDRDIAKEESYSTISLLSTDTKILMSKYK
		·				QLKSSDL
581	1931	Α	4414	670	3	VLVHRQCGGILRLRRKEAVSVLDSADIEVTDS
		1				RLPHATIVDHRPQHRWLETCNAPPQLIQGKA
						RSAPKPSQASGHFSVELVRGYAGFGLTLGGG
[		l				RDVAGDTPLAVRGLLKDGP\AQRCGRLEVGD
						LVLHINGESTQGLT\HAQAVERIRAGGPQLHL
						VIRRPLETHPGKPRGVGEPRKGVVPSWPDRSP
						DPGGPEVTGSRSSSTSLVQHPPSRTTLKKTRG
500	1000		1404	101	1.10	SPE
582	1932	Α	4424	194	449	VLYIRKKKRLEKLRHQLMPMYNFDPTEEQDE
						LEQELLEHGRDAASVQAATSVQAMQGKTTL
583	1933	A	4435	1	166	PS\QGPLQRPSRLVFT\DVANAIHV
363	1933	A	4433	1	100	APGPPVPPPGSPPEQMPGPCPASMPP/DPPPGS PPEQMPGPCPVSAPP/GPPPGSPPEQMPGPCPV
						SAPPALLQDTSV
584	1934	A	4439	1	628	SATPQQPSAPQHQGTLNQPPVPGMDESMSYQ
364	1754	^	4437	1	028	APPQQLPSAQPPQPSNPPHGAHTLNSGPQPGT
						APATQHSQAGPATGQAYGPHTYTEPAKPKK
						GQQLWNRMKPAPGT\EV\$SST\$R\$DPLLLPPR
						ALAPTQRASTVVLAPSPT/SEKVQNHSGSSAR
						GNLSGKPDDWP/LGHERVCGALLHRL*VGGG
		ļ				QGPHGKAAQGGAAGAAAGRLGLYH
585	1935	A	4463	10	144	HKPVTNSRDTQEVPLEKAKQVLKIIATFKHTT
						SIFDDFAHYEKRQ
586	1936	A	4464	1309	103	LNAESYVSFTTKLDIPTAAKYEYGVPLQTSDS
						FLRFPSSLTSSLCTDNNPAAFLVNQAVKCTRK
		ļ				INLEQCEEIEALSMAFYSSPEILRVPDSRKKVPI
						TVQSIVIQSLNKTLTRREDTDVLQPTLVNAGH
						FSLCVNVVLEVKYSLTYTDAGEVTKADLSFV
						LGTVSSVVVPLQQKFEIHFLQENTQPVPLSGN
		1	1			PGYVVGLPLAAGFQPHKGSGIIQTTNRYGQLT
						ILHSTTEQDCLALEGVRTPVLFGYTMQSGCK
						LRLTGALPCQLVAQKVKSLLWGQGFPDYVA
						PFGNSQGP/ADMLDWVPIHFITQSFNRKDSCQ
						LPGALVIEVKWTKYGSLLNPQAKIVNVTANLI
						SSSFPEANSGNERTILISTAVTFVDVSAPAEAG
	1000					FRAPPAINARLPFNFFFPFV
587	1937	A	4471	614	387	LLGRASAC/LQLQSSW/D/HRPMLPYLANFVF
			J			CKDR/SFTWLPRLVLNSWLQVILLPWPPTGCD
500	1022		4400	1000	1450	NKHEPPCPATKRRHSGSI
- VV 1	1938	A	4480	1720	1458	HDLGSLQPPPPGFKRFSCLSLPSSWDYRLMPP
588	1730	i e				CPANFCIIII/DFLVETGFHHVGQASHELLTSGD
200	1730					
						PPTSASQSAGITGMSYHTWFGES
589	1939	Ā	4487	922	332	PPTSASQSAGITGMSYHTWFGÈS APVTTSPRVGQPW/RTALALRSLYRARPSLRC
		A	4487	922	332	PPTSASQSAGITGMSYHTWFGÈS APVTTSPRVGQPW/RTALALRSLYRARPSLRC PPVELPWAPRRGHRLSPADDELYQRTRISLLQ
		A	4487	922	332	PPTSASQSAGITGMSYHTWFGÈS APVTTSPRVGQPW/RTALALRSLYRARPSLRC

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						DTPNACATFNFLCHEGRVTGAALIPPPGGTSL TSLGQAAQ
590	1940	A	4492	1	472	FFFFETESRSVAQAGVQWRDLGSLQAPPPGFT PFSCLSLPSSWDYRRPPLRPANFFVFLVETGFP RFSRDGLDLLT/S/GDPPTSASQSAGITGVSHR ARPKRIGEPRRKCGNAVVWPSTSLGDHRVTS VPHQGGLPGPIRVAPSSAGQREASQGPPGR
591	1941	A	4495	1444	1116	IAARFTLAKTWNQLKRP\TMIDSIKKTR\YIYT MEYYADTERNEIMSF\AGTWVELEAIILSKLM LKDNWVEDTIPQGAVPCTATAEGMKRLLFAL EPWDSSCFPHPSSGV
592	1942	A	4496	2	919	RTRPLFSGRPTRPVCTMSDERRLPGSAVGWL VCGGLSLLANAWGILSVGAKQKKWKPLEFL LCTLAATHMLNVAVPIATYSVVQLRRQRPDF EWNEGLCKVFVSTFYTLTLATCFSVTSLSYHR MWMVCWPVNYRLSNAKKQAGHTVMGIWM GSFILSALPAVGWHDTSERFYTHGCRFIVAEI GLGFGVCFLLLVGGSVAMGVICTAIALFQTL AVQVGRQADHRAFTVPTIVVEDAQGKRRSSI DGSEPAKTSLQTTGLVTTIVFIYDCLMGFPVL GPFSLADTHLSDLPYTWGDRDSGGACVM
593	1943	A	4506	2	193	FFFEAESCSVPQAGVQRPDLGWLHAPPP\GSC HFPASASQVAGTTHARHHTQLIF\AFLVENGL C
594	1944	A	4507	1327	647	KMAGGVRPLRGLRALCRVLLFLSQFCILSGG ESTEIPPYVMKCPSNGLCSRLPADCIDCTTNFS CTYGKPVTFDCAVKPSVTCVDQDFKSQKNFII NMTCRFCWQLPETDYECTNSTSCMTVSCPRQ RYPANCTVR\DHVHCLGNRTFPKMLYCNWT GGYKWVYGLWLLRHHPRWGLGADRF\YLGP VAGTASGKLFSFGGLGIWTLIDVLLIGVGYVG PADGSLYI
595	1945	A	4512	533	264	FFFKMESYSVARLECSGAISAPCNLHLLGSNN SPASASRV/AGNIGARHHTQQIFVLLVQMRVH YVGQDGLDLL/NLMIHPPRSPKVLGLQA
596	1946	A	4513	3	1674	HASDHLYPNFLVNELILKQKQRFEEKRFKLD HSVSTNGHRWQIFQDWLGTDQDNLDLANV NLMLELLVQKKKQLEAESHAAQLQILMEFLK VARRNKREQLEQIQKELSVLEEDIKRVEEMS GLYSPVSEDSTVPQFEAPSPSHSSIIDSTEYSQP PGFSGSSQTKKQPWYNSTLASRRKRLTAHFE DLEQCYFSTRMSRISDDSRTASQLDEFQEC\LS KF\TRYNSVRPL\ATLSYASDLYNGSQYKSLV FEFDRDCDYFAIAGVTKKIKVYEYDTVIQDA VDIHYPENEMTCNSKISCISWSSYHKNLLASS DYEGTVILWDGFTGQRSKVYQEHEKRCWSV DFNLMDPKLLASGSDDAKVKLWSTNLDNSV ASIEAKANVCCVKFSPSSRYHLAFGCADHCV HYYDLRNTKQPIMVFKGHRKAVSYAKFVSG EEIVSASTDSQLKLWNVGKP\YCLRSFKGHIN EKNFVIGLASNGDYIACGSENNSLYLYYKGLS KTLLTFKFDTVKSVLDKDRKEDDTNEFVSAV CWRALPDGESNVLIAANS\QGTI\KVLELV
597	1947	Ā	4518	536	824	RSLALSPGLECSGMISAHCNLHLLGSSDPPTS ASQVAEITSVRHHTWLIFCI\LGQMGFHHVGE
598	1948	A	4524	1	384	QAGLELLTSWDPAILPSQSAGIIGMSPHAWPP FDTEFVNIGGDFDAAAGVFR\CRLPGAYFFSF TLGKLPRKTLSVKLMKNRDEVQAMIYDDGSS

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						YGAYSNHGKYITFSGFLVYPDLAPAAPPGLG ASELL
599	1949	A	4526	366	776	MGQPAPYAEGPIQGGDAGELCKCDFLVFTSP NPEAVCEAGTPAMFQTAWRQMESCSI/AQAG VQWRDPGSLHPPPLGFKRFSCLSLPSSWDYK HAPPHPANFCIFSRDQVSPCWPGWSRSLDLVI PPPWLPKVLGLQA
600	1950	A	4529	776	334	FFFETESCYVAQAGVQWCDLCSLQAPPPG\SS DPPASASRVAGTTGARHHTQLIFVFLVETGFH \MLARDGLKLLTSSDPPASASQSSWDYRREPP RLANFFVFLVETGSRYVAQAGVQWLFTGAIP LLISTGVLTCSVSDLGRFTPP
601	1951	A	4533	1460	403	HEVQESIHFLESEFSRGISDNYTLALITYALSS VGSPKAKEALNMLTWRAEQEGGMQFWVSSE SKLSDSWQPRSLDIEVAAYALLSHFLQFQTSE GIPIMRWLSRQRNSLGGFASTQDTTVALKALS EFAALMNTERTNIQVTVTGPSSPSPVKFLIDT HNRLLLQTAELADGTANGSV/SISANGFGFAI CQLNVVYNVKASGSSRRRRSIQNQEAFDLDV AVKENKDDLNHVDLNVCTSFSGPGRSGMAL MEVNLLSGFMVPSEAISLSETVKKVEYDHGK LNLYLDSVNETQFCVNIPAVRNFKVSNTQDA SVSIVDYYEPRRQAVRSYNSEVKLSSCDLCSD VQRLPSL
602	1952	A	4540	1963	295	MRAPGRPALRPLPLPPLLLLLSSPWGRAVPC VSGGLPKPANITFLSINMKNVLQWTPPEGLQG VKVTYTVQYFIYGQKKWLNKSECRNINRTYC DLSAETSDYEHQYYAKVKAIWGTKCSKWAE SGRFYPFLETQIGPPEVALTTDEKSISVVLTAP EKWKRNPEDLPVSMQQIYSNLKYNVSVLNT KSNRTWSQCVTNHTLVLTW\LEPNTLYCVHV ESFVPGPPRRAQPSEKQCARTLKDQSSEFKAK IIFWYVLPISITVFLFSVMGYSIYRYIHVG\KEK HP\ANLILIYG\NEFDKRFFVPA\EKIV\INFI\TL NIS\DDSKISHQDMSLLGKSSDVSSLNDPQPSG NLRPPQEEEEVKHLGYASHLMEIFCDSEENT\ EGTSFTQQESLSRTIPPDKTVIEYEYDVRTTDI CAGPEEQELSLQEEVSTQGTLLESQAALAVL GPQTLQYSYTPQLQDLDPLAQEHTDSEEGPEE EPSTTLVDWDPQTGRLCIPSLSSFDQDSEGCE PSEGDGLGEEGLLSRLYEEPAPDRPPGENETY LMQFMEEWGLYVQMEN
603	.1953	A	4543	3	600	YSAVEFVEQASGISDWWNPALRKRMLSDSGL GMIAPYYEDSDLKDLSHSRVLQSPVSSEDHAI LQAVIAGDLMKLIESYKNGGSLLIQGPDHCSL LHYAAETGNGEIVKYILDHGPSELLDMADSE TGETALHKAACQRNRAVCQLLVDAGASLRK\ TDSKGKTPQERAQQA\GDPDLAA/YTIESRQN YKVIGHEDLETAV
604	1954	A	4548	3	938	QDNKVQNGSLHQKDTVHDNDFEPYLTGQAN QSNSYPSMSDPYLSSYYPPSIGFPYSLNEAPW STAGDPPIPYLTTYGQLSNGDHHFMHDAVFG QPGGLGNNIYQHRFNFFPENPAFSAWGTSGS QGQQTQSSAYGSSYTYPPSSLGGTVVDGQPG FHSDTLSKAPGMNSLEQGMVGLKIGDVSSSA VKTVGSVVSSVALTGVLSGNGGTNVNMPVS KPTSWAAIASKPAKPQPKMKTKSGPVMGGG LPPPPIKHNMDIGTWDNKGPVPKAPVPQQAP

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605	1955	A	4553	2	2304	QPPQ ILLQEKRNCLLMQLEEATRLTSYLQSQLKSLC
			4555		2504	ASTLTVSSGSSRGSLASSRGSLASSRGSLSSVS FTDIYGLPQYEKPDAEGSQLLRFDLIPFDSLGR DAPFSEPPGPSGFHKQRRSLDTPQSLASLSSRS SLSSLSPPSSPLDTPFLPASRDSPLAQLADSCE GPGLGALDRLRAHASAMGDEDLPGMAALQP HGVPGDGEGPHERGPPPASAPVGGTVTLRED SAKRLERRARRISACLSDYSLASDSGVFEPLT KRNEDAEEPAYGDTASNGDPQIHVGLLRDSG SECLLVHVLQLKNPAGLAVKEDCKVHIRVYL PPLDSGTPNTYCSKALEFQVPLVFNEVFRIPV HSSALTLKSLQLYVCSVTPQLQEELLGIAQIN LADYDSLSEMQLRWHSVQVFTSLNHQGRGR LGVQERAPPGTLHTPSPSPA/STDAVTVLLAR TTAQLQAVERELAEERAKLEYTEEEVLEMER KEEQAEAISERSWQADSVDSGCSNCTQTSPPY PEPCCMGIDSILGHPFAAQAGPYSPEKFQPSPL KVDKETNTEDLFLEEAASLVKERPSRRARGSP FVRSGTIVRSQTFSPGARSQYVCRLYRSDSDS STLPRKSPFVRNTLERRTLRYKQSCRSSLAEL MARTSLDLELDLQASRTRQRQLNEELCALRE LRQRLEDAQLRGQTDLPPWVLRDERLRGLLR EAERQTRQTKLDYRHEQAAEKMLKKASKEI YQLRGQSHKEPIQVQTFREKIAFFTRPRINIPPL PADDV
606	1956	A	4555	3429	776	PGSGPGPAPFLAPVAAPVGGISFHLQIGLSREP VLLLQDSSGDYSLAHVREMACSIVDQKFPEC GFYGMYDKILLFRHDPTSENILQLVKAASDIQ EGDLIEVVLSASATFEDFQIRPHALFVHSYRA PAFCDHCGEMLWGLVKQGLKCEGCGLNYH KRCAFKIPNNCSGVRRRLSNVSLTGVSTIRT SSAELSTSAPDEPLLQKSPSESFIGREKRSNSQ SYIGRPIHLDKILMSKVKVPHTFVIHSYTRPTV CQYCKKLLKGLFRQGLQCKDCRFNCHKRCA PKVPNNCLGEVTINGDLLSPGAESDVVMEEG SDDNDSERNSGLMDDMEEAMVQDAEMAMA ECQNDSGEMQDPDPDHEDANRTISPSTSNNIP LMRVVQSVKHTKRKSSTVMKEGWMVHYTS KDTLRKRHYWRLDSKCITLFQNDTGSRYYKE IPLSEILSLEPVKTSALIPNGANPHCFEITTANV VYYVGENVVNPSSPSPNNSVLTSGVGADVAR MWEIAIQHALMPVIPKGSSVGTGTNLHRDISV SISVSNCQIQENVDISTVYQIFPDEVLGSGQFGI VYGGKHRKTGRDVAIKIIDKLRFPTKQESQLR NEVAILQNLHHPGVVNLECMFETPERVFVVM EKLHGDMLEMILSSEKGRLPEHITKFLITQILV ALRHLHFKNIVHCDLKPENVLLASADPFPQV KLCDFGFARIIGEKSFRRSVVGTPAYLAPEVL RNKGYNRSLDMWSVGVIIYVSLSGTFPFNED EDIHDQIQNAAFMYPPNPWKEISHEAIDLINN LLQVKMRKRYSVDKTLSHPWLQDYQTWLDL RELECKIGERYTTHESDDLRWEKYAGEQGLQ YPTHLINPSASHSDTPETEETEMKALGERVSIL
607	1957	A	4563	1	4499	SRPWWLRASERPSAPSAMAKRSRGPGRRCLL
						ALVLFCAWGTLAVVAQKPGAGCPSRCLCFRT TVRCMHLLLEAVPAVAPQTSILDLRFNRIREI QPGAFRRLRNLNTLLLNNNQIKRIPSGAFEDL ENLKYLYLYKNEIQSIDRQAFKGLASLEQLYL

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uence			914	ng to first amino acid residue of peptide sequence	acid residue of peptide sequence	Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
						HFNQIETLDPDSFQHLPKLERLFLHNNRITHL VPGTFNHLESMKRLRLDSNTLHCDCEILWLA DLLKTYAESGNAQAAAICEYPRRIQGRSVATI TPEELNCERPRITSEPQDADVTSGNTVYFTCR AEGNPKPEIIWLRNNNELSMKTDSRLNLLDD GTLMIQNTQETDQGIYQCMAKNVAGEVKTQ EVTLRYFGSPARPTFVIQPQNTEVLVGESVIL ECSATGHPPPRISWTRGDRTPLPVDPRVNITPS GGLYIQNVVQGDSGEYACSATNNIDSVHATA FIIVQALPQFTVTPQDRVVIEGQTVDFQCEAK GNPPPVIAWTKGGSQLSVDRRHLVLSSGTLRI SGVALHDQGQYECQAVNIIGSQKVVAHLTVQ PRVTPVFASIPSDTTVEVGANVQLPCSSQGEP EPAITWNKDGVQVTESGKFHISPEGFLTINDV GFADAGRYECVARNTIGSASVSMVLSVNVPD VSRNGDPFVATSIVEAIATVDRAINSTRTHLF DSRPRSPNDLLALFRYPRDPYTVEQARAGEIF ERTLQLIQEHVQHGLMVDLNGTSYHYNDLVS PQYLNLIANLSGCTAHRRVNNCSDMCFHQKY RTHDGTCNNLQHPMWGASLTAFERLLKSVY ENGFNTPRGINPHRLYNGHALPMPRLVSTTLI GTETVTPDEQFTHMLMQWGQFLDHDLDSTV VALSQARFSDGQHCSNVCSNDPPCFSVMIPPN DSRARSGARCMFFVRSSPVCGSGMTSLLMNS VYPREQINQLTSYIDASNVYGSTEHEARSIRD LASHRGLLRQGIVQRSGKPLLPFATGPPTECM RDENESPIPCFLAGDHRANEQLGLTSMHTLW FREHNRIATELLKLNPHWDGDTIYYETRKIVG AEIQHITYQHWLPKILGEVGMRTLGEYHGYD PGINAGIFNAFAT\AAFRFGHTLVNPLLLPGLD ENFQPIAQDHLPLHKAFFSPFRIVNEGGIDPLL RGLFGVAGKMRVPSQLLNTELTERLFSMAHT VALDLAAINIQRGRDHGIPPYHDYRVYCNLS AAHTFEDLKNEIKNPEIREKLKRLYGSTLNID LFPALVVEDLVPGSRLGPTLMCLLSTQFKRLR DGDRLWYENPGVFSPAQLTQIKQTSLARILCD NADNITRVQSDVFRVAEFPHGYGSCDEIPRVD LRVWQDCCEDCRTRGQFNAFSYHFRGRRSLE FSYQEDKPTKKTRPRKIPSVGRQGEHLSNSTS A\FSTRSDASG\TNDFQRVCSWEMQKTITDLR TQIKKLESRLSTTECVDAGGESHANNTKWK KDACTICECKDGQVTCFVEACPPATCAVPVNI PGACCPVCLQKRAEEKP
608	1958	A	4566		1135	FSFLC/GVSGRLGLDSEEDYYTPQKVDVPKAL IIVAVQCGCDGTFLLTQSGKVLACGLNEFNKL GLNQCMSGIINHEAYHEVPYTTSFTLAKQLSF YKIRTIAPGKTHTAAIDERGRLLTFGCNKCGQ LGVGNYKKRLGINLLGGPLGGKQVIRVSCGD EFTIAATDDNHIFAWGNGGNGRLAMTPTERP HGSDICTSWPRPIFGSLHHVPDLSCRGWHTILI VEKVLNSKTIRSNSSGLSIGTVFQSSSPGGGGE GGPDAW
609	1959	A	4567	P. C. C. C. C. C. C. C. C. C. C. C. C. C.	412	FFFFETESRSVAQAGVQWRDLGSLQAPPPGFT PFSCLSLPSSWDYRRPPLRPANFFVFLVETGF HRFSRDGLDLLT/S/GDPPASASQSAGITGVSH RARPRINLRNVIYSFAVTYCLNYISLAMSSTL KLSFHVLSGS
610	1960	A	4570	697	467	ECRGVISAH\CCTLCLPSSSDSASAF\RVARTT GTCDYAQLIFAFLVEMGFHHVGQDGLHLL/N LVIRPPRPPKVLGLQA

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first	Predicted end nucleotide .location corresponding to last amino acid residue	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
				amino acid residue of peptide sequence	of peptide sequence	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
611	1961	A	4571	25	1396	ADPHTTVIRFFPAASATKRVLPPVLRVSSPRT WNPNVPESPRIPAPRLPKRMSGAPTAGAALM LCAATAVLLSAQGGPVQSKSPRFASWDEMN VLAHGLLQLGQG\CANT\GAHPQSAERAGA\R LSACGSACQGTEGSTDLPLAPESRVDPEVLHS LQTQLKAQNSRIQQLFHKVAQQQRHLEKQHL RIQHLQSQFGLLDHKHLDHEVAKPARRKRLP EMAQPVDPAHNVSRLHRLPRDCQELFQVGER QSGLFEIQPQGSPPFLVNCKMTSDGGWTVIQR RHDGSVDFNRPWEAYKAGFGDPHGEFWLGL EKVHSITGDRNSRLAVQLRDWDGNAELLQFS VHLGGEDTAYSLQLTAPVAGQLGATTVPPSG LSVPFSTWDQDHDLRRDKNCAKSLSGGWWF GTCSHSNLNGQYFRSIPQQRQKLKKGIFWKT WRGRYYPLQATTMLIQPMAAEAAS
612	1962	A	4575	162	3	FFFETESRSVAQAGVQWRDLSSLQPPPPG\SR GSPASASPVAGITGTRHHRTRG
613	1963	A	4584	687	321	PLAQRRPFLWVTVKTNGHIWGSSTYPHFWGS SNS/PASASQVAGIPNARHQARIIFVFLVEPRF HHVGRAGLGFL/NLAICLPQHPKVLGLQACN LNIKPHPAHKYISMIQFNVHFMCMSVHIYI
614	1964	A	4589	727	299	PGSAQSAQRGRGRRRARAGSATQITMYSFMG GGLFCAWVGTILLVVAMATDHWMQYRLSGS FAHQGLWRYCLGNKCYLQTDSIAYWNATRA FMILSALCAISGIIMGIMAF/GWVAVLMTFFA GIFYMCAYRVHECRRLSTPR
615	1965	A	4590	2	414	TILPEKIQAWAQKQCPQSGEEAVALVVHLEK ETGRLRQQVSSPVHREKHSPLGAAWEVADFQ PEQVETQPRAVSREEPGSLHSGHQEQLNRKR ERRPLPKNARPSPWVPALADEWNTLHQEVTT TRLPAGSQEPVKD
616	1966	A	4592	773	488	DFALVAQAGVQWHNLGSPQPLPPGFKRFSCL SLPSSWEYRCVPP/RLANFVFLVEMGFLHVGQ AGLELPTSGDPPALASQSAGITGVTTVPSGPG
617	1967	В	4595	84	478	XRHGLREPLLERRCAAASSFQHSSSLGRELPY DPVDTEGFGEGGDMQERFLFPEYILDPEPQPT REKQLQELQQQQEEEERQRQQRREERRQQNL RARSREHPVVGHPDPALPPSGVNCSGCGAEL HCQDAR*
618	1968	A	4596	2945	1188	ARSRNSARGVYGMCVDTLFLCFLEDLERNDG SAERPYFMCSTLKKPLARRCFPAIHAYKGVL MVGNETTYEDGHGSRKNITDLVEGAKKANG VLEARQLAMRIFEDYTVSWYWIIIGLVIAMA MSLLSIILLHLLAGIMGWVMIIMEISELGYRIF HCYMEYSRLRGEAGSDVSLVDLGFQTDFRV YLHLRQTWLAFMIILSILEVIIILLIFLRKRILI AIALIKEASRAVGYVMCSLLYPLVTFFLLCLCI AYWASTAVFLSTSNEAVYKIFDDSPCPFTAKT CNPETFPSSNESRQCPNARCQFAFYGGESGYH RALLGLQIFNAFMFFWLANFVLALGQVTLAG AFASYYWALRKPDDLPAFPLFSAFGRALRYH TGSLAFGALILAIVQIIRVILEYLDQRLKAAEN KFAKCLMTCLKCCFWCLEKFIKFLNRNAYIM IAIYGTNFCTSARNAFFLLMRNIIRVAVLDKV TDFLFLLGKLLIVGSVGILAFFFTHRIRIVQDT APPLNYYWVPILTVIVGSYLIAHGFFSVYGMC VDTLFLCFLEDLERNDGSAERPYFMSSTLKKL LNKTNKKAAES
619	1969	A	4601	2	357	RTSVEPYILGEF/RKLSNNTKVVKTEYKATEY

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide scquence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  GLAYGHFSYEFSNHRDVVVDLQGWVTGNGK GLIYLTDPQIHSVDQKVFTTNFGKRGIFYFFN
620	1970	A	4606	1	2415	NQHVECNEICHRLSLTRPSMEKPCKS  MERLWGLFQRAQQLSPRSSQTVYQRVEGPR KGHLEEEEEDGEEGAETLAHFCPMELRGPEP LGSRPRQPNLIPWAAAGRRAAPYLVLTALLIF TGAFLLGYVAFRGSCQACGDSVLVVSEDVN YEPDLDFHQGRLYWSDLQAMFLQFLGEGRL EDTIRQTSLRERVAGSAGMAALTQDIRAALS RQKLDHVWTDTHYVGLQFPDPAHPNTLHWV DEAGKVGEQLPLEDPDVYCPYSAIGNVTGEL VYAHYGRPEDLQDLRARGVDPVGRLLLVRV GVISFAQKVTNAQDFGAQGVLIYPEPADFSQ DPPKPSLSSQQAVYGHVHLGTGDPYTPGFPSF NQTQFPPVASSGLPSIPAQPISADIASRLLRKL KGPVAPQEWQGSLLGSPYHLGPGPRLRLVVN NHRTSTPINNIFGCIEGRSEPDHYVVIGAQRDA WGPGAAKSAVGTAILLELVRTFSSMVSNGFR PRRSLLFISWDGGDFGSVGSTEWLEGYLSVL HLKAVVYVSLDNAVLGDDKFHAKTSPLLTSL IESVLKQVDSPNHSGQTLYEQVVFTNIPSWD\ AEVIRPLPM\DSSAY\SFTAFVGVPAVEFSFME\ DDQ\AYPFLHTKEDTYENLHKVLQGRLPAVA QAVAQLAGQLLIRLSHDRLLPLDFGRYGDVV LRHIGNLNEFSGDLKARGLTLQWVYSARGDY IRAAEKLRQEIYSSEERDERLTRMYNVRIMRV EFYFLSQYVSPADSPFRHIFMGRGDHTLGALL DHLRLLRSNSSGTPGATSSTGFQ\ESRFRRQL\
621	1971	A	4610	793	334	ALL\TWDACKGAANALSGDVWNIDNNF ISRVDDFVGSGIANVIIAVAIFSIPAFARLVRG\ NTLVLKQQTFIESARSIGASDMTVLLRHILPGT GSSIVVFFTMRIGTSIISAASLSFLGLGAQPPTP EWGAMLNEARADMVIAPHVAVFPALAIFLTV LAFNLLGDGLRDALDPKIKG
622	1972	A	4614	2	820	LVYVMIAIFCIASAMSLYNCLAALIHKIPYGQ CTIACRGKNMEVRLIFLSGLCIAVAVVWAVF RNEDRWAWILQDILGIAFCLNLIKTLKLPNFK SCVILLGLLLLYDVFFVFITPFITKNGESIMVEL AAGPFGNNEKNDGNLVEATGQPSAPHEKLPV VIRVPKLIYFSVMSVCLMPVSILGFGDIIVPGL LIAYCRRFDVQTGSSYIYYVSV\TVAYAIGMIL TFVVLG\LMKKGQPALLYLVPCTLITA/CQFV AWETVREMKKFWERVTS
623	1973	A	4619	17	691	TLVSVVEFVRRADLTREDLAPSSVDSGQAGF GGCCESGLPNTMPSAFSVSSFPVSIPAVLTQT DWTEPWLMGLATFHALCVLLTCLSSRSYRLQ IGHFLCLVILVYCAEYINEAAAMNWRLFSKY QYFDSRGMFISIVFSAPLLVNAMIIVVMWVW KTLNVMTDLKNAQERRKEKKRRKED*GAA AAWSLRPSRPPSAAPSAAVCVAWASFQLTHG LKNRCFI
624	1974	Α .	4622	164	668	VSCYTALQSIMNQPESANDPEPLCAVCGQAH SLEENHFYSYPEEVDDDLICHICLQALLDPLD TPCGHTYCTLCLTNFLVEKDFCPMDRKPLVL QHCKKSSILVNKLLNKLLVTCPFREHCTQVL QRCDLEHHFQTSQAWGTHL*SQLLGRLRQED CLSPGVHHCSEV
625	1975	A	4625	474	473	CFLSPSPLLPPLLLSSSSSPSFPLPPPPTLLPSTLP PPLLIPSS*LSP

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
626	1976	A	4629	249	3	KLKGNECFCYHCNVCIFLMIKK*GLFLC*IYFI LFFET*SHSFTRLECSGTISAHCSLQLQGSSNSP ASASQVAGIAGTHH
627	1977	A	4635	1	301	FFFFETKPFFAPQAGGQGPSRGSLNPLPTGLK QFSGLTLSRSGNNGPRPPPRVNFGILRGNGVP PGGAG*PRPPDLRGPPGLAPPQGGNNGGDPP ARAYL
628	1978	A	4648	1357	782	KLFSSQRLFGPHIQAINPSFLLLSFFPS*LLAMR TVGNNAFILVFLVYRIVLLLF*HV*PAYFQPSK NKTAKINCN*RPFLFLVCYLL*AELHIGIFIANF YDCIPNKLNEHLWPKLLQSLIFHVDFCGFLHK VFYICFTEFLLFLYFL*LFIIKVSCSII*CSTICVF SYKSFAVIIFFVDNTRFFSFGF
629	1979	A	4660	18	999	HHELHTLELLQNPKEVLTRSEIQDVNYSLEAV KVKTVCQIPLMKEMLKRFQVAVNLAEDTAH PKLVFSQEGRYVKNTASASSWPVFSSAWNYF AGWRNPQKTAFVERFQHLSCVLGKNVFTSG KHYWEVESRDSLEVAVGVCREDVMGITDRS KMSPDVGIWAIYWSAAGYWPLIGFPGTPTQQ EPALHRVGVYLDRGTGNVSFYSAVDGVHLH TFSCSSVSRLRPFFWLSPLASLVIPPVTDRK*G FSSPDQNSFPVVQLRDTHPWALFCPSCLYPG WSIFWVSLTVPFGICPLCASQEAVPWEVGLA NGDGTGNFPRRFWEIFL
630	1980	A	4669	2	358	FFFFFETESHSVAQAGMQWRNLGSLPAPPPGF TPFFCLSLLNGWDYRRPPPHLANFFVLLVETG FHDVGQDGLDLLTS*STPSASQSAEITGVSHC TRLKKIRFAKGHVEFFFESHVE
631	1981	A	4674	953	614	TPIRGTDDEHEECTVQEYSAGKNTCLRPGAV AHTCNPCTLGGRGRWIT*GSGVQDQPGPTWQ NPVFLERRPRALHSSPGLTTQRILWAQGLWV GAGSTGCSRGPRGEGVFREG
632	1982	A	4678	34	314	RSTHASGMISPSFGFMGHLLRLEFEILPSTPNP *LPSYQGEAAGSSLISHLQTFSPDLKGVYCTFP ASGLAPVPTHWTVSELSRSPVATATFC
633	1983	A	4696	1	1365	RTLGMEGERRASQAPSSGLPAGGANGESPGG GAPFPGSSGSSALLQAEVLDLDEDEDDLEVFS KDASLMDMNSFSPMMPTSPLSMINQIKFEDEP DLKDLFITVDEPESHVTTIETFITYRITTKTSRG EFDSSEFEVRRRYQDFLWLKGKLEEAHPTLII PPLPEKFIVKGMVERFNDDFIETRRKALHKFL NRIADHPTLTFNEDFKIFLTAQAWELSSHKQ GPGLLSRMGQTVRAVASSMRGVKNRPEEFM EMNNFIELFSQKINLIDKISQRIYKEEREYFDE MKEYGPIHILWSASEEDLVDTLKDVASCIDRC CKATEKRMSGLSEALLPVVHEYVLYSEMLM GVMKRRDQIQAELDSKVEVLTYKKADTDLL PEEIGKLEDKVECANNALKADWERWKQNM QNDIKLAFTDMAEENIHYYEQCLATWESFLT SQTNLHLEEASEDKP
634	1984	A	4708	421	158	SYWVGEDYTYKFFEVILIDPFHKAIRRNPDTQ WISKAVYKHREMCGLTSTGRKSHGLEKDRM FPHAIGGSCRAA*RRKTLQFPCYH
635	1985	A	4709	42	341	YIKQPDAKERRRTVHWKKETESEASEITIPPST PGVPQAPGHWEDYGRGDNFYLPH*DPGGIVL WNIFNRMPIARKNITDGEHHEYLIEVPRLFHT SED
636	1986	A	4721	2	351	EKPDHFFPEGTSFIHEPRRPN*GDLVHCLGGIS RSTTVTVA*LMQKLNLSMNDAYYIVIMKMSS

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SEQ ID	SEQ ID	Met hod	SEQ ID NO:	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of nucl-	NO: of peptide	поа	in NO:	beginning nucleotide	nucleotide location	D=Aspartic Acid, E=Glutamic Acid,
eotide	seq-		USSN	location	corresponding	F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
	uence		914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
uence		[	914	amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of		Y=Tyrosine, X=Unknown, *=Stop codon,
					sequence	/=possible nucleotide deletion, \=possible
1		1		peptide		
		ļ		sequence		nucleotide insertion ISPNFNSMDQPLDFQRTLGLRSPCYNRVPAQK
637	1007		1770	664	0.50	MYFTTPSNHNAYQVDSVQST
03/	1987	A	4726	004	253	NTGLTCSIQRKCGETQLYRREENRLILLLQDH
						LKSESFQVLTLSPRLEFSGLISAHCNLRLPGSS
						DSSASSRAAGITGVHHHAWLIFFFLVETGFL
						HAG*AGLELLTSGDPPASASRSAGITGVSHHA
(20	1988	ļ <del>.</del>	477.4	0.4	500	RPRETRFL
638	1988	A	4734	24	592	GGMDSRVSGTTSNGETKPVYPVMEKKEEDG
						TLERGHWNNKMEFVLSVAGEIIGLGNVWRFP
						YLCYKNGGGAFFIPYLVFLFTCGIPVFLLETAL
						GQYTSQGGVTAWRKICPIFEGIGYASQMIVIL
		Į				LNVYYIIVLAWALFYLFSSFTIDLPWGGCYHE
						WNTEHCMEFQKTNGSLNGTSENATSPVIEFW
639	1989	A	4743	1040	699	QGLTLLPRMECSATITAHCSLELPGSIDLPTSA
						S*VARTTGTHHHPWLILVLLL*TWGSYYVAQ
1	1	ł	ł	}		AGLELLGSSNLPAAMVSQSAQIIGHDHCAWA
						TSNHVLYTQEGLRRGKEG
640	1990	Α	4771	527	2	GRIDCPHPATVLAQPIFIDACSVLGAYQGAQN
						WIRRRPCLPSGCLKMNREIGPLQHSLCCPGWS
						QTPGLKAILLRQPPK*LGLQMESHSCPPAWSA
		ĺ	1			MARSRLTATSASQVQAILLPQPPGTTDSCSPS
						PDHEQQPLSWVLPPPQKDMNPREQQVALGP
						QAAALPWAVWRNDCFPR
641	1991	A	4780	16	473	RPSSQCGGIPTGWKKGLAPELSSELSSPPLPAR
	]	ļ	J			LQLAASPYFSPSWAECPQPVPAGTHATWCLA
						RVWARMTPPGPAGIPSHPLPPPPPERSVPIPSP
		1				FPARDSGSRQGHSTDRYKHTDAPRDAHRRVP
						QRDTDTGVHTGSGTHTHAHTPPEK
642	1992	A	4798	1	487	GYSFRCDIVDYSRSPTALRMARTCWLYYFSK
		!	}			FIELLDTIFFVLRKKNSQVTFLHVFHHTIMPW
						TWWFGVKFAAGGLGTFHALLNTAVHVVMY
						SYYGLSALGPAYQKYLWWKKYLTSLQLVQF
J .						VIVAIHISQFFFMEDCKYQFPVFACIIMSYSFM
						FLLLFLH
643	1993	A	4799	2	391	LMAFIEMHISGSLVYLKIKTKIYSYFSMLNFLL
						QEIPLSEILRISSPRDFTNISQGSNPHCFEIITDT
						MVYFVGENNGDSSHNPVLAATGVGLDVAQS
1		ł	1	}		WEKAIRQALMPVTPQASVCTSPGQGKDHSK
						Q*ASVCTSPGQGKDHSKQ
644	1994	A	4800	488	101	AYPLFAVHPVHTECVAGVVGRAYLLCALFFL
						LSFLGYCKAFRESNKEGAHSSTFWVLLSIFLG
						AVAMLCKEQGITVLVRAATWLGPAFSVCPFP
[			1			SYKDIWGWPCLCGVLHAYIPLLV
645	1995	A	4805	458	126	LLWTTVLCQTPARPQSTMIHLGHILFLLLLPV
]						AAAQTTPGERSSLPAFYPGTSGSCSGCGSLSL
	1	1	l		}	PLLAGLVAADAVASLLIVGAVFLCARPRRSP
						AQEDGKVYINMPGRG
646	1996	A	4817	47	1033	LQGDTWHLSFLSHFSRLHGGVPGRGLLEGNL
"	1,,,0	11	1017	**	1055	LQPQAPGHDMTSIPFPGDRLLQVDGVILCGLT
[		[	ĺ			HKQAVQCLKGPGQVARLVLERRVPRSTQQC PSANDSMGDERTAVSLVTALPGRPSSCVSVT
1						
						DGPKF*SSN*KRIANGLGFSFVQMEKESCSHL
		İ				KSDLVRIKRLFPGHPAEENGAIAAGDIILGRE
] ]			1			WEGPRKASSSRCRGSWAMQLSVQAGPSFAS
						YYPAAVEVLHLLRGAPQEVTLLLCRPPPGAL  DELEGENOTEEL CARVETT ATCTROCTORU
Į						PELEQEWQTPELSADKEFTRATCTDSCTSPIL
						GSRGQLGGTVPPQMQGKAWGLRPESSQKAIR
647	1007	_	1051	1044	225	EGTMGAKTERDLGPVP
1 104 / 1	1997	I A	4854	1044	335	PRVRGDWPLEKKKSNSNIHPIFSWCGSTDSKD

000 00	Lorge	136:	Long	1 5 11	1	
SEQ ID NO: of	SEQ ID NO: of	Met	SEQ ID NO:	Predicted	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide	nou	in in	beginning nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence			914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
402.00				amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
				peptide		/=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion
						IVMPTYDLTDSVLETMGRVSLDMMSVQANT
1	1					GPPWESKNSTAVWRGRDSRKERLELVKLSRK
						HPELIDAAFTNFFFFKHDENLYGPIVKHISFFD
		1				FFKHKYQINIDGTVAAYRLPYLLVGDSVVLK
						QDSIYYEHFYNELQPWKHYIPVKSNLSDLLEK
	Ì	ľ	1			LKWAKDHDEEAKKIAKAGQEFARNNLMGD
648	1998	<u> </u>	4867	2030	837	DIFCYYFQTFPRNMPIYK
648	1998	Α	4867	2030	837	AGMLPAVGSADEEEDPAEEDCPELVPMETTQ
	1					SEEEEKSGLGAKIPVTIITGYLGAGKTTLLNYI
						LTEQHSKRVAVILNEFGEGSALEKSLAVSQG GELYEEWLELRNGCLCCSVKDNGLRAIENLM
			1			QKKGKFDYILLETTGLADPGAVASMFWVDA
						ELGSDIYLDGIITIVDSKYGLKHLAEEKPDGLI
						NEATRQVALADAILINKTDLVPEEDVKKLRT
						TIRSINGLGQILETQRSRVDLSNVLDLHAFDSL
1		l		}	l	SGISLQKKLQHVPGTQPHLDQSIVTITFDVPG
						NAKEEHLNMFIQNLLWEKNVRNKDNHCMEV
		l				IRLKGLVSIKDKSQQVIVQGVHELYDLEETPV
						SWKDDTERTNRLVLLGRNLDKDILKQLFIAT
						VTETEKQWTTHFKEDQVCT
649	1999	A	4873	226	189	DGVSLLLPKLGVQWAQYWAHWQPPLPGFKR
						FSCLSLRSSWD*KCAPPHPAFVFLVEMGFHRV
						GQAGLELRTSGDPPASASQSAGITGVSHLA*P
650	2000	- A	4874	-	120	TSMPLLPFQRLCVYI
630	2000	A	4874	2	437	FFFLRRSFAFVAQAGVQWCDLGSPQPLPPGF K*FSCLSLPSSWDYRHAPPPCPS*FLYF**RQG
						FTMLARLVLNS*PHDLPTSPSQSAEIKGVSHR
						CPASFYLFLKYYLEAKFCA*GECAPSAGVGA
ĺ		İ				GYKRGHKSCLLINCVVQI
651	2001	A	4898	1701	771	DAWGPETRLARILNPDSFIEPRPGRLPELEATR
				-7		PHMEPKASCPAAAPLMERKFHVLVGVTGSV
						AALKLPLLVSKLLDIPGLEVAVVTTERAKHFY
,						SPQDIPVTLYSDADEWEMWKSRSDPVLHIDL
						RRWADLLLVAPLDANTLGKVASGICDNLLTC
	ļ	ļ				VMRAWDRSKPLLFCPAMNTAMWEHPITAQQ
						VDQLKAFGYVEIPCVAKKLVCGDEGLGAMA
						EVGTIVDKVKEVLFQHSGFQQS*PGISVMGVP
						LYSEWVQAKSVKMDVGKIGGYPHLLNGGPA
652	2002	_	4927	1	611	LSLPRGQACSRLNWTEGPGLSFFQPGEAAA
0.5.2	2002	A	4921	1	011	FRGRQTSRPARGFSPWRPPGTMQEPSSGECPA
			1			SP*LPCASNRLAFGGLIFPCAPLVPYPAPFSPLL PAFSCAPRPRAHTHSRTHPSAPLVPKPSSRAR
						GQSPIPSRASSPSCSWAQVPGVALARCAGVC
		}	1	1	}	KPGDSWRVAACISGRCCSRGRRRGSGPRNPE
						QSFRGAWGPSFWGSWKSQRELSAGGAQAWP
						LLGSAGSGLRGEA
653	2003	A	4965	2	283	FFFFI*DGVSLCHPGWNAVARSWLTATSASR
				_		VQAVSCFRLPSSWDYRHATMPG*FF*YF**R
						WGFTILAILVLNS*PQVICPPWPPKVLTLQA
654	2004	A	4968	3 .	437	RPGIPGRRFRRSWFCQLP*EPEPGLESLATPGD
						IPAVGLGALGVIPPVRVPQRPPTQRSQGRGW
						DPERDPGCRVQVSRGPRFGEQKTPGLQGCLP
						PPCLTHLAAASCVVVWCGRWKRDSAECQCD
						HSCSAVSQQEDRCRSSSCS
655	2005	A.	4983	201	397	MNNNTTCIQPSMISSMALPIIYILLCIVGVFGN
						TLSQWIFLTKIGKKTSTHIYLSHLVTANLLVC
656	2006	A	4988	332	159	LVHKDMYREFFEEEAQASNKHVTRCLTSLVI
						REVHIKTMR*HFLPIRLEKNKNNIKD
657	2007	В	5008	129	465	MAGMKTASGDYIDSSWELRVFVGEEDPEAES

000 75	1 one to	1 3 6 4	1.000	T 20 10 11	I 70 10 10 10 10 10 10 10 10 10 10 10 10 10	
SEQ ID NO: of	SEQ ID NO: of	Met hod	SEQ ID NO:	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
nucl-	peptide	noa	in in	beginning nucleotide	nucleotide location	D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence	ĺ	09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence	denoe		914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
		İ	7	amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
!		ļ	1.	peptide		/=possible nucleotide deletion, \=possible
		1		sequence		nucleotide insertion
						VTLRVTGESHIGGVLLKIVEQINRKQDWSDH
						AIWWEQKRQWLLQTHWTLDKYGILADARLF
						FGPQHRPVILRLPNRRALRLX*
658	2008	A	5017	1	292	FFFFKETESHSVTQAGVQWHDLGSLQPPPPGF
						KRFSCLSLLSSWDYRCAPPHPANFVFLVETGF
						HHVAQAGLKLLTL*SANLGLSTSLPIPLFILLS
659	2009	A	5018	17	338	RGHGGKSLTGGTPGNWGDGLLVSEDWSHLIF
						T*NSLVSPVLGKWSPCLQGPGLSAVHTWPWL
		ļ		<u> </u>	ļ	MAACWAVHVKTHMRPGLAVLPRLVLNSWS
	0010		5000		010	*AIILLWPPKALGLQA
660	2010	Α	5028	2	310	SRVDDFVGERRGGCDECLCGHRGLRAVPLG
		1				HPGHLCLQPPGGPA*FLDYCRGCCPHPVPGST
	i	1				AGSCPRQKKTTPGPTVLCVCSFWIYQRGEPH
661	2011	A	5050	752	431	HRTGARWNH
001	2011	Α.	3030	132	451	RQSCSSTQAKVQWFHYGPLQSQPPGLKQSSQ LSLPNSRDHRHVPPRLAIFSFAETGSPYFAQAS
						LELLGSSHPPTSASQSARITGVSHRAWPLK*F
	]				ļ	NLNQYQTLTMN
662	2012	A	5054	48	103	ELNNGPFQMPLCNGGNLAVTGSWADRSPLH
002	2012	**	3031	10	103	EAASQGRLLALRTLLSQGYNVNAVTLDHVTP
						LHEACLGDHVACARTLLEAGANVNAITIDGV
	ł	ł				TPLFNACSQGSPSCAELLLEYGAQAQLESCLP
						SPTHEGASKGHHECLDILISWGIDVDQEIPHSG
						TPLYVACMAQQFHCIWNLIYAGAGVRKGKY
		]				WDTPLPGAGHQSTQKLE*LFAMVEIWQ
663	2013	A	5066	951	580	VRNS*SFAHCASVYKHHYMDGQTPCLFVSSK
						ADLPEGVAVSGPSPAEFCRKHRLPAPVPFSCA
						GPAEPSTTIFTQLATMAAFPHLVHAELHPSSF
		İ				WLRGLLGVVGAAVAAVLSFSLYRVLVKSQ
664	2014	Α	5071	550	1	LSFIEVLSMEQVNKTVVREFVVLGFSSLARLQ
						QLLFVIFLLLYLFTLGTNAIIISTIVLDRALHTP
	1		1	ļ		MYFFLAILSCSEICYTFVIVPKMLVDLLSQKK
						TISFLGCAIQMFSFLFFGSSHSFLLAAMGYDR
						YMAICNPLRYSVLMGHGVCMGLMAAAWAC
	-			10.6		GFTVSLVTTSLVFHLPFHSSNQHE
665	2015	A	5074	496	692	QQYHNTGSAGHHAHCQVGHSPHVHYPSGCG
		[				PL*IQRGLPSFNSLEGHSLKDSGHEESVQLDSE
		}				HDVQRSLYCDTAVNDVLNTSVTSMGSQMPD
						HDQNEGFHCREECRILGHSDRCWMPRNPMPI
		]		]		RSKSPEHVRNIIALSIEATAADVEAYDDCGPT KRTFATFGKDVSDHPAEERPTLKGKRTVDVT
						ICSPKVNSVIREAGNGCEAISPVTSPLHLKSSL
						PTKPSVSYEIVDPGITARRC
666	2016	A	5080	408	248	IMLLSTSS*VYFQSSTKDSHFFLFDFQKTGPPL
JUU	2010	l ''	3000	100	270	VGPKAQLSGLQLQPCLYKRR
667	2017	A	5081	129	247	DLTNSHFFLFDFQKTGPPLGGPKAQFSSLQLQ
		1	5551	1		PCVY*RR
668	2018	A	5086	852	233	NIKSNDRWVQIKTAYKYFF*KNGDNYNWVF
		l · •	5555	30		RALPTTFADIENLKYLLFTRDASQPFYLGHTV
						IFGDLEYVTVEGGIVLSRELMKRLNRLLDNSE
		l				TCADQSVIWKLSEDKQLAICLKYAGVHAENA
		[		[		EDYEGRDVFNTKPIAQLIEEALSNNPQQVVEG
						CCSDMAITFNGLTPQKMEVMMYGLYRLRAF
		Ì				GHYFNDTLVFLPPVGSEND
669	2019	Α	5101	1	329	PGRPTRPPLLTLLAHVSPEPAGPSCDSLAQPG
		i				ASGV*VQHDSHPPLLCGSQCLSEPVPGSHGPP
				l l		Tibe : QIIDBIM : EECOSQCEEDEI VI COITGII
						RGCQHEAAPCPRGPGSDGLHHASAACASLPP
670	2020	A	5102	3	547	

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  DEL*NMNGRVDYLVTEEEINLTRGPSGLGFNI
						VGGTDQQYVSNDSGIYVSRIKENGAAALDGR LQEGDKILSVNGQDLKNLLHQDAVDLFRNA GYAVSLRVQHRLQVQNGPIGHRGEGDPSGIPI FMVLVPVFALTMVAAWAFMRYRQQL
671	2021	A	5105	672	400	RDGREELCLQQEPTLPSRICSSAPLLYFLFICPF VLLLLLISLLCLYWKARKLSTLRSNTRKEKA LWVDLKEAGGVTTNRMED*EEDECN
672	2022	A	5148	72	314	IIYFSYNIFLKITELLNDVERLKQALNGLSQLT YTSGNPTKRQSQLIDTLQHQVKSLEQQLAVS NQAHGALQEYVLAPCS
673	2023	A	5152	210	335	REILCSRIGRLNIV*MSLFPNLTCRLNAIPIKIPA NHFVEVT
674		A	5153	3	2953	LTEDQPFDILQKSLQEANITEQTLAEEAYLDA SIGSSQQFAQAQLHPSSSASFTQASNVSNYSG QTLQPIGVTHVPVGASFASNTVGVQHGFMQH VGISVPSQHLSNSSQISGSGQIQLIGSFGNHPS MMTINNLDGSQIILKGSGQQAPSNVSGGLLV HRQTPNGNSLFGNSSSSPVAQPVTVPFNSTNF QTSLPVHNIIIQRGLAPNSNKVPINIQPKPIQM GQQNTYNVNNLGIQQHHVQQGISFASASSPQ GSVVGPHMSVNIVNQQNTRKPVTSQAVSSTG GSIVIHSPMGQPHAPQSQFLIPTSLSVSSNSVH HVQTINGQLLQTQPSQLISGQVASEHVMLNR NSSNMLRTNQPYTGPMLNNQNTAVHLVSGQ TFAASGSPVIANHASPQLVGGQMPLQQASPT VLHLSPGQSSVSQGRPGFATMPSVTSMSGPSR FPAVSSASTAHPSLGSAVQSGSSGSNFTGDQL TQPNRTPVPVSVSHRLPVSSSKSTSFSNTPGT GTQQGFFCQAQKKCLNQTSPISAPKTTDGLR QAQIPGLLSTTLPGQDSGSKVISASLGTAQPQ QEKVVGSSPGHPAVQVESHSGGVRPAAKQ LTKGAFILQQLQRDQAHTVTPDKSHFRSLSD AVQRLLSYHVCQGSMPTEEDLRKVDNEFETV ATQLLKRTQAMLNKYRCLLLEDAMRINPPAE MVMIDRMFNQEERASLSRDKRLALVDPEGFQ ADFCCSFKLDKAAHETQFGRSDQHGSKASSS LQPPAKAQGRDRAKTGVTEPMNHDQFHLVP NHIVVSAEGNISKKTECLGRALKFDKVGLVQ YQSTSEEKASRREPLKASQCSPGPEGHRKTSS RSDHGTESKLSSILADSHLEMTCNNSFQDKSL RNSPKNEVLHTDIMKGSGEPQPDLQLTKSLET TFKNILELKKAGRQPQSDPTVSGSVELDFPNF SPMASQENCLEKFIPDHSEGVVETDSILEAAV NSILEC
675	2025	A	5154	599	1880	LKKMEPFSCDTFVALPPATVDNRIIFGKNSDR LYDEVQEVVYFPAVVHDNLGERLKCTYIEID QVPETYAVVLSRPAWLWGAEMGANEHGVCI GNEAVWGREEVCDEEALLGMDLVRLGLERA DTAEKALNVIVDLLEKYGQGGNCTEGRMVF SYHNSFLIADRNEAWILETAGKYWAAEKVQE GVRNISNQLSITTKIAREHPDMRNYAKRKGW WDGKKEFDFAAAYSYLDTAKMMTSSGRYCE GYKLLNKHKGNITFETMMEILRDKPSGINME GEFLTTASMVFILPQDSSLPCIHFFTGTPDPER SVFKPFIFVPHISQLLDTSSPTFELEDLVKKKS HFKPDRRHPLYQKHQQALEVVNNNEEKAKI MLDNMRKLEKELFREMESILQNKHLDVEKIV NLFPQCTKDEIQIYQSNLSVKVSS

CEOID	L CEO ID	N/o+	020	Dun dinte d	D	I A district C C di
SEQ ID NO: of	SEQ ID NO: of	Met hod	SEQ ID NO:	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide	liou	in NO.	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence			914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
				amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
		-		residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
1		}		peptide	1 .	/=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion
676	2026	Α	5155	2	306	FFFLRRSLALSPRPDCGLQWRNLGSLQAPPPG
						FTPFSCLSLPSSWDYRRPPPRPANFLYF**RRG
1						FTLLARMVSIS*PHDPPASASQSAGITGVSHRA
						RPT
677	2027	Α	5167	97	740	FFHSVDLLALEQSKTFYKPDWFDIVESEVKCC
						KEAVCVIDMSSFTEFEITSTGDQALEVLQYLF
						SNDLDVPVGHIVHTGMLNEGGGYENDCSIAR
						LNKRSFFMISPTDQQVHCWAWLKKHMPKDS
1	Ì	ľ	1		ĺ	NLLLEDVTWKYTALNLIGPRAVDVLSELSYA
						PMTPDHFPSLFCKEMSVGYANGIRVMSMTHT
						GEPGFMLYIPIEYRWGFTMLSTLVSNS
678	2028	A	5183	1919	2018	PALCRLRDDMTVCVADFGLSKKIYSGDYYRQ
		İ			1	GRIAKMPVKWIAIESLADRVYTSKSDVWAFG
	ĺ	ì	1	1		VTMWEIATRGMTPYPGVQNHEMYDYLLHG
						HRLKQPEDCLDELCKI**SPQSP
679	2029	Α	5190	39	499	RESQVKHFKMRKIDLCLSSEGSEVILATSSDE
			1	-		KHPPENIIDGNPETFWTTTGMFPQEFIICFHKH
						VRIERLVIQSYFVQTLKIEKSTSKEPVDFEQWI
	1					EKDLVHTEGQLQNEEIVAHDGSATYLRFIIVS
						AFDHFASVHSVSAEGTVVSNLSS
680	2030	Α	5204	541	92	EILAVLKLACGDISLNALALMVATAVLTLAPL
						LLICLSYLFILSAILRVPSAAGRCKAFSTCSAH
						RTVVVVFYGTISFMYFKPKAKDPNVDKTVAL
						FYGVVTPSLNPIIYSLRNAEVKAAVLTLLRGG
(0)	0001			10		LLSRKASHCYCCPLPLSAGIG
681	2031	A	5207	10	247	VPDNGDVTKLPVCSTLVEETSLTVSEAMEQSI
						KNESPLPGTLAHTCNTSTLGGRGRWIT*GREF
700	2022				001	DTSMANMVKPCLYRK
682	2032	A	5210	2	231	FFFETESYSITQAGVQWPNLSSLKTLPPGFK*F
l		l	ł			SCLSLPSSWDYRCLPPCPANFCIFSRNGVLPC
692	2033		5212	05	102	WPGWSRTPDLS
683	2033	A	5218	85	402	CPSVSGLIKSDLRRHNINIGITNVDVKAVSNIF
						MILLRSMYRINVKPYFFI*LFFSRVNC*SVIIG
						YARCYTFLIF*LFL*IPADSPTDQEPKTVMLSK QSESAI
684	2034	A	5220	1	194	NLMKEMONLNSENHKTWEEYKDTK*IMSYF
004	2034	Α.	3220	,	194	YG*ALNVIKMAVLPKLMYRFSATLVKIPQHL
						TDS
685	2035	A	5228	260	440	LHSQDGNSDPRKPQGEMSAHAFPVQTCGEED
005	2033	Α	3226	200	140	QKKTPQVPINFTELSKCS*S*KIMSGERE
686	2036	A	5239	79	508	GGEAAARAAKLSSPRPHRVGRRERGVGGMS
000	2030	A	3237	17	300	AFSEAALEKKLSELSNSQQSVQTLSLWLIHHR
						KHSRPIVTVWERELRKAKPNRKLTFLYLAND
						VIQNSKRKGPEFTKDFAPVIVEAFKHVSSETD
						ESCKKHLGRVLSIWEERS
687	2037	A	5244	1	428	MAAVVAATALKGRGARNARVLRGILAGATA
007	2031	<sup>43</sup>	3244	1	720	NKASHNRTRALQSHSSPEGKEEPEPLSPELEYI
						PRKRGKNPMKAVGLAWAIGFPCGILLFILTKR
						EVDKDRVKQMKARQNMRLSNTGEYESQRFR
						ASSOSAPSPDVGSGVOT
688	2038	A	5249	1	1407	LQQTEDKSLLNQGSSSEEVAGSSQKMGQPGP
1	2000	1 1 2	32-47	1	1-107	SGDSDLATALHRLSLRRQNYLSEKQFFAEEW
						QRKIQVLADQKEGVSGCVTPTESLASLCTTQS
						EITDLSSASCLRGFMPEKLQIVKPLEGSQTLY
	,					HWQQLAQPNLGTILDPRPGVITKGFTQLPGD
,			,			AIYHISDLEEDEEEGITFQVQQPLEVEEKLSTS
		ľ		<b>!</b>		KPVTGIFLPPITSAGGPVTVATANPGKCLSCT
						NSTFTFTTCRILHPSDITQVTPSSGFPSLSCGSS
						GSSSNTAVNSPALAYRLSIGESITNRRDSTTT
				L		

SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO; of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine.
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence	J	j	914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
				amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
ĺ	[		ĺ	residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon.
ł			ł	peptide	1	/=possible nucleotide deletion, \=possible
ļ				sequence		nucleotide insertion
	J				]	FSSTMSLAKLLQERGISAKVYHSPISENPLQPL
						PKSLAIPSTPPNSPSHSPCPSPLPFEPRVHLSEN
						FLASRPAETFLQEMYGLRPSRNPPDVGQLKM
	1	1	1	}		NLVDRLKRLGIARVVKNPGAQENGRCQEAEI
	1					GPQKPDSAVYLNSGSSLLGGLRRNQSLPVIM
689	2039	$+$ $\overline{A}$	5254	12	2621	GSFAAPVCTSSPKMGVLKED
002	2035	11	3234	2	2021	LSLFGSRALGRSGARAMAKAKKVGARRKAS
}	ł	1	1			GAPAGARGGPAKANSNPFEVKVNRQKFQILG
	J		j	]		RKTRHDVGLPGVSRARALRKRTQTLLKEYKE
						RDKSNVFRDKRFGEYNSNMSPEEKMMKRFA LEQQRHHEKKSIYNLNEDEELTHYGQSLADIE
	1	ĺ	ĺ	ĺ	ĺ	KHNDIVDSDSDAEDRGTLSGELTAAHFGGGG
						GLLHKKTQQEGEREKPKSRKELIEELIAKSK
'}		1		}	ļ	QEKRERQAQREDALELTEKLDQDWKEIQTLL
1						SHKTPKSENRDKKEKPKPDAYDMMVRELGF
1	1				Ì	EMKAQPSNRMKTEAELAKEEQEHLRKLEAE
1	}	1	1	1	ł	RLRRMLGKDEDENVKKPKHMSADDLNDGFV
1	}	ļ				LDKDDRRLLSYKDGKMNVEEDVQEEQSKEA
						SDPESNEEEGDSSGGEDTEESDSPDSHLDLES
į.	1	1				NVESEEENEKPAKEQROTPGKGLISGKERAG
1	ł		1			KATRDELPYTFAAPESYEELRSLLLGRSMEEQ
						LLVVERIQKCNHPSLAEGNKAKLEKLFGFLLE
						YVGDLATDDPPDLTVIDKLVVHLYHLCOMFP
ſ	ſ		[			ESASDAIKFVLRDAMHEMEEMIETKGRAALP
1	ľ		1			GLDVLIYLKITGLLFPTSDFWHPVVTPALVCL
		ļ	1 .			SQLLTKCPILSLQDVVKGLFVCCLFLEYVALS
		]				QRFIPELINFLLGILYIATPNKASQGSTLVHPFR
					•	ALGKNSELLVVSAREDVATWQQSSLSLRWA
1	1			ľ		SRLRAPTSTEANHIRLSCLAVGLALLKRCVLM
1						YGSLPSFHAIMGPLRALLTDHLADCSHPQELQ
		1				ELCQSTLTEMESQKQLCRPLTCEKSKPVPLKL
						FTPRLVKVLEFGRKQGSSKEEQERKRLIHKHK
Í	1					REFKGAVREIRKDNOFLARMQLSEIMERDAE
690	2040	A	5261	1 .	304	RKRKVKQLFNSLATQEGEWKALKRKKFKK
					JVT	FFFFVFLVETGFHHVGQAGLELLTSGDPPTW ASQSAGITGVSHCSWPVIYVLSTLLHAVRNVL
		[	ſ		[	FKRTFPLKSSSFLSYDKEIFPILIVLKFYLVTLT
					1	SFVK
691	2041	A	5270	3	158	NCHTTHCTANWVHLPGTPPGWKIDGPAAAL
				1		EVLSSFFFFFLKFSYKPQNIV
692	2042	A	5282	56	1268	GMEPVGCCGECRGSSVDPRSTFVLSNLAEVV
						ERVLTFLPAKALLRVACVCRLWRECVRRVLR
1		1	ł	ł		THRSVTWISAGLAEAGHLEGHCLVRVVAEEL
		J		ļ		ENVRILPHTVLYMADSETFISLEECRGHKRAR
[	]	J		J	J	KRTSMETALALEKLFPKQCQVLGIVTPGIVVT
						PMGSGSNRPQEIEIGESGFALLFPQIEGIKIOPF
1		1	1	1		HFIKDPKNLTLERHQLTEVGLLDNPELRVVLV
		}			1	FGYNCCKVGASNYLQQVVSTFSDMNIILAGG
		ł	1		1	QVDNLSSLTSEKNPLDIDASGVVGLSFSGHRI
					1	QSATVLLNEDVSDEKTAEAAMQRLKAANIPE
	1	- 1	- 1		İ	HNTIGFMFACVGRGFQYYRAKGNVEADAFR
			1	ļ	1	KFFPSVPLFGFFGNGEIGCDRIVTGNFILRKCN
693	2042	<del></del>	5201			EVKDDDLFHSYTTIMALIHLGSSK
ひどろ	2043	A	5301	362	507	EEIKERFGPGLVIYWYGFIQELDCNRERGILLK
694	2044		5210		204	ACFPTNIVTLCHSIA
U74	ZU44	A	5310	1	204	RVLTAINHTLKENLRKFYKGKKDKPLDLRPK
				ļ		KTRAMRRLNMHEENLKTKKQHRKERLYPL
695	2045	A	5315	125	1506	RKYAAKA
	20-13	4.3	2213	143	1596	ETRSTAVKSEVQVCISLLLCLEDRTMPKKAKP

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
						TGSGKEEGPAPCKQMKLEAAGGPSALNFDSP SSLFESLISPIKTETFFKEFWEQKPLLIQRDDPA LATYYGSLFKLTDLKSLCSRGMYYGRDVNV CRCVNGKKKVLNKDGKAHFLQLRKDFDQKR ATIQFHQPQRFKDELWRIQEKLECYFGSLVGS NVYITPAGSQGLPPHYDDVEVFILQLEGEKH WRLYHPTVPLAREYSVEAEERIGRPVHEFML KPGDLLYFPRGTIHQADTPAGLAHSTHVTIST YQNNSWGDFLLDTISGLVFDTAKEDVELRTG IPRQLLLQVESTTVATRRLSGFLRTLADRLEG TKELLSSDMKKDFIMHRLPPYSAGDGAELSTP GGKLPRLDSVVRLQFKDHIVLTVLPDQDQSD ETQEKMVYIYHSLKNSRETHMMGNEEETEFH GLRFPLSHLDALKQIWNSPAISVKDLKLTTDE EKESLVLSLWTECLIQVV
696	2046	A	5318	1476	742	LMKXYLEAAELGEISDIHTKLLRLSSSQGTIET SLQDIDSRLSPGGSLADAWAHQEGTHPKDRN VEKLQVLLNCMTEIYYQFKKDKAERRLAYN EEQIHKFDKQKLYYHATKAMTHFTDECVKK YEAFLNKSEEWIRKMLHLRKQLLSLTNQCFDI EEEVSKYQEYTNELQETLPQKMFTASSGIKHT MTPIYPSSNTLVEMTLGMKKLKEEMEGVVKE LAENNHILESGGSLTMDGGLRNVDCL
697	2047	A	5320	244	478	LDYNFFLFEMTFGLVSQAGVQWHDLGSLQPP PPGFKQFSCLSLPSSWDYRHLPPHLANFSREG VSPSWPGWSRTPDFR
698	2048	A	5324	266	714	LPIRKSLRSVRSGFPTSQSPITRNLDGTASGSC LAKTVTGSLFRINVGLRGLVAGGIIGALLGTP VGGLLMAFQKYSGETVQERKQKDRKALHEL KLEEWKGRLQVTEHLPEKIESSLQEDEPENDA KKIEALLNLPRNPSVIDKQDKD
699	2049	Α	5334	699	277	RPHGHLVCISSSAGLSGVNGLADYCASKFAA FGFAESVFVETFVQKQKGIKTTIVCPFFIKTGM FEGCTTGCPSLLPILEPKYAVEKIVEAILQEKM YLYMPKLLYFMMFLKSFLPLKTGLLIADYLGI LHAMDGFADOKK
700	2050	A	5344	3	614	PTAEEMSSLTPESSPELAKRSWFGNFISLDKEE QIFLVLKDKPLSSIKADIVHAFLSIPSLSHSVLS QTSFRAEYKASGGPSVFQKPVRFQVDISSSEG PEPSPRRDGSGGGGIYSVTFTLISGPSRRFKRV VETIQAQLLSTHDQPSVQALADEKNGAQTRP AGAPPRSLQPPPGRPDPELSSSPRRGPPKDKK LLATNGTPL
701	2051	A	5346	3	1383	HASVLFCRVMAASKTQGAVARMQEDRDGSC STVGGVGYGDSKDCILEPLSLPESPGGTTTLE GSPSVPCIFCEEHFPVAEQDKLLKHMIIEHKIV IADVKLVADFQRYILYWRKRFTEQPITDFCSV IRINSTAPFEEQENYFLLCDVLPEDRILREELQ KQRLREILEQQQQERNDTNFHGVCMFCNEEF LGNRSVILNHMAREHAFNIGLPDNIVNCNEFL CTLQKKLDNLQCLYCEKTFRDKNTLKDHMR KKQHRKINPKNREYDRFYVINYLELGKSWEE VQLEDDRELLDHQEDDWSDWEEHPASAVCL FCEKQAETIEKLYVHMEDAHEFDLLKIKSELG LNFYQQVKLVNFIRRQVHQCRCYGCHVKFKS KADLRTHMEETKHTSLLPDRKTWDQLEYYFP TYENDTLLWTLSDSESDLTAQEQNENVPIISE DTSKLYALKOSSILNOLLI
702	2052	A	5356	2502	1540	DTSKLYALKQSSILNQLLL MAAATRGCRPWGSLLGLLGLVSAAAAAWD

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703	2053	A	5380	278	657	LFLQKLRMKTEEEARTHTEIEMFLRKEQQKL EERLEFWMEKYDKDTEMKQNELNALKATKA SDLAHLQDLAKMIREYEQVIIEDRIEKERSKK KVKQDLLELKSVIKLQAWWRGTMIRREIGGF KM
704	2054	A	5381	1	1003	FRGRAVKMAAVVEVEVGGGAAGERELDEV DMSDLSPEEQWRVEHARMHAKHRGHEAMH AEMVLILIATLVVAQLLLVQWKQRHPRSYN MVTLFQMWVVPLYFTVKLHWWRFLVIWILF SAVTAFVTFRATRKPLVQTTPRLVYKWFLLIY KISYATGIVGYMAVMFTLFGLNLLFKIKPEDA MDFGISLLFYGLYYGVLERDFAEMCADYMA STIGFYSESGMPTKHLSDSVCAVCGQQIFVDV SEEGIIENTYRLSCNHVFHEFCIRGWCIVGKK QTCPYCKEKVDLKRMFSNPWERPHVMYGQL LDWLRYLVAWQPVIIGVVQGINYILGLE
705	2055	A	5396	3	675	IYDRDPLQLATRAGQPLDINMAGEPKPYRPKP GNKRPLSALYRLESKEPFLSVGGYVFDYDYY RDDFYNRLFDYHGRVPPPPRAVIPLKRPRVA VTTTRRGKGVFSMKGGSRSTASGSTGSKLKS DELQTIKKELTQIKTKIDSVLGRLDKIEKQQK AEAEAQKKLLEESLVLIQEECVSEIADHSTEEP AEGGPDADGEEMTDGIEEAFDEDGGHELFLQ IK
706	2056	A	5410	2	98	GRVGLNLEGRGCSEPKWRHCTPTWATEQDSI S
707	2057	A	5415	6	287	PFKLTPSFLSHAFSSGQERKVFIELNHIKKCNT VRGVFVLEEFGNYTILLLGLDSHGSNSNLGAP EEGLGAGRKRTSVEKSGGAGVTRKKRDP
708	2058	Ā	5423	3	291	SSSNPLGSPSTLWKLCSFVLHNKSCCCSFFGS TPTLRAITLTVRVCGFIPEVSKTTNPLGRTNNS GCTIFKTVTLTARSTASLLKSVRPRTHQKE
709	2059	Α	5424	679	347	RIRHEEKRGSRGRGRRTSEEDTPKKKKHKGG SEFTDTILSVHPSDVLDMPVDPNEPTYCLCHQ VSYGEMIGCDNPDCPIEWFHFACVDLTTKPK GKWFCPRCVQEKRKKK
710	2060	A	5442	1073	559	QESLKKKIQPKLSLTLSSSVSRGNVSTPPRHSS GSLTPPVTPPITPSSSFRSSTPTGSEYDEEEVDY EESDSDESWTTESAISSEAILSSMCMNGGEEK PFACPVPGCKKRYKNVNGIKYHAKNGHRTQI RVRKPFKCRCGKSYKTAQGLRHHTINFHPPV SAEIIRKMQQ
711	2061	A	5449	1	319	GDSLCVPQYNKYREERVILFLKMASGHAFQP DLVKRIRDAIRMGLSARHVPSLILETKGIPYTL NGKKVEVAVKQIIAGKAVEQGGAFSNPETLD LYRDIPELQGF
712	2062	A	5499	91	749	RPTPGHGDFWMQPLTKDAGMSLSSVTLASAL QVRGEALSEEEIWSLLFLAAEQLLEDLRNDSS DYVVCPWSALLSAAGSLSFQGRVSHIEAAPF

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						KAPELLQGQSEDEQPDASQMHVYSLGMTLY WSAGFHVPPHQPLQLCEPLHSILLTMCEDQPH RRCTLQSVLEACRVHEKEVSVYPAPAGLHIR RLVGLVLGTISEVSREPCFSSSSCWSCVAIKI
713	2063	A	5506	22	478	VEELILVSRLDPHLHTPMYFFLAHLSFLDLSFT TSSIPQLLYNLNGCDKTISYMGCAIQLFLFLGL GGVECLLLAVMAYDRCVAICKPLHYMVIMN PRLCRGLVSVTWGCGVANSLAMSPVTLRLPR CGHHEVDHFLCEMPALIRMACISTV
714	2064	A	5514	25	220	AIRPYWCENNIIGIGKLSTADGKAFADPEVLR RLTSSVSCALDEAAAALTRMRAESTANAGQS DK
715	2065	A	5526	3	810	KVTAPRRPQRYSSGHGSDNSSVLSGELPPAM GRTALFHHSGGSSGYESLRRDSEATGSASSAP DSMSESGAASPGARTRSLKSPKKRATGLQRR RLIPAPLPDTTALGRKPSLPGQWVDLPPPLAG SLKEPFEIKVYEIDDVERLQRPRPTPREAPTQG LACVSTRLRLAERRQQRLREVQAKHKHLCEE LAETQGRLMLEPGRWLEQFEVDPELEPESAE YLAALERATAALEQCVNLCKAHVMMVTCFD ISVAASAAIPGPQEVDV
716	2066	A	5529	458	790	SPGYGENKFTVTSXNIAVPLCEMNKIYSYYSD SSSSERTMDLVLEMCNTNSIHWCGISGRQLG KLHPSSSLCLALTLLSSVQGLQSISGLRLTDTF LKRTYEYDDIAQVCV
717	2067	A	5531	3	460	NSEDLLKYFNPESWOEDLDNMYLDTPRYRG RSYHDRKSKVDLDRLNDDAKRYSCTPRNYS VNIREELKLANVVFFPRCLLVQRCGGNCGCG TVNWRSCTCNSGKTVKKYHEVLQFEPGHIKR RGRAKTMALVDIQLDHHERCDCICSSRPPR
718	2068	A	5586	311	88	AVLKNMAPMTALGLLDLHILNLILFLSAGEDF TSVVSEIMMYILLVFLTLWLLIEMIYCYRKVS KAEEAAQENA
719	2069	A	5598	1	330	KNCANEAVVQKILDRVLSRYDVRLRPNFGSM LATNSTRGLNEDELMAHGQEKDSSSESEDSC PPSPGCSFTEGFSFDLLNPDYVPKVDKWSRFL FPLAFGLFNIVAAERC
720	2070	A .	5628	798	148	LPPAQIPEAWLLLANVVVVLILVPLKDRLIDP LLLRCKLLPSALQKMALGMFFGFTSVIVAGV LEMERLHYIHHNETVSQQIGEVLYNAAPLSIW WQIPQYLLIGISEIFASIPGLEFAYSEAPRSMQG AIMGIFFCLSGVGSLLGSSLVALLSLPGGWLH CPKDFGNINNCRMDLYFFLLAGIQAVTALLF VWIAGRYERASQGPASHSRFSRDRG
721	2071	A	5632	146	536	MSALIVRKLRSAELTLFSELPTVLGANVNAA KLHETALHHAAKVKNVDLIEMLIEFGGNIYA RDNRGKKPSDYTWSSSAPAKCFEYYEKTPLT LSQLCRVNLRKATGVRGLEKIAKLNIPPRLID YLSYN
722	2072		5638	3	3806	CPSLDIRSEVAELRQLENCSVVEGHLQILLMF TATGEDFRGLSFPRLTQVTDYLLLFRVYGLES LRDLFPNLAVIRGTRLFLGYALVIFEMPHLRD VALPALGAVLRGAVRVEKNQELCHLSTIDW GLLQPAPGANHIVGNKLGEECADVCPGVLGA AGEPCAKTTFSGHTDYRCWTSSHCQRVCPCP HGMACTARGECCHTECLGGCSQPEDPRACV ACRHLYFQGACLWACPPGTYQYESWRCVTA ERCASLHSVPGRASTFGIHQGSCLAQCPSGFT RNSSSIFCHKCEGLCPKECKVGTKTIDSIQAA

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				·		QDLVGCTHVEGSLILNLRQGYNLEPQLQHSL GLVETITGFLKIKHSFALVSLGFFKNLKLIRGD AMVDGNYTLYVLDNQNLQQLGSWVAAGLTI PVGKIYFAFNPRLCLEHIYRLEEVTGTRGRQN KAEINPRINGDRAACQTRTLRFVSNVTEADRI LLRWERYEPLEARDLLSFIVYYKESPFQNATE HVGPDACGTQSWNLLDVELPLSRTQEPGVTL ASLKPWTQYAVFVRAITLTTEEDSPHQGAQS- PIVYLRTLPAAPTVPQDVISTSNSSHLLVRW KPPTQRNGNLTYYLVLWQRLAEDGDLYLND YCHRGLRLPTSNNDPRFDGEDGDPEAEMESD CCPCQHPPPGQVLPPLEAQEASFQKKFENFLH NAITIPISPWKVTSINKSPQRDSGRHRRAAGPL RLGGNSSDFEIQEDKVPRERAVLSGLRHFTEY RIDIHACNHAAHTVGCSAATFVFARTMPHRE ADGIPGKVAWEASSKNSVLLRWLEPPDPNGL ILKYEIKYRRLGEEATVLCVSRLRYAKFGGV HLALLPPGNYSARVRATSLAGNGSWTDSVAF YILGPEEDAGGLHVLLTATPVGLTLLIVLAA LGFFYGKKRNRTLYASVNPEYFSASDMYVPD EWEVPREQISIIRELGQGSFGMVYEGLARGLE AGEESTPVALKTVNELASPRECIEFLKEASVM KAFKCHHVVRLLGVVSQGQPTLVIMELMTR GDLKSHLRSLRPEAENNPGLPQPALGEMIQM AGEIADGMAYLAANKFVHRDLAARNCMVSQ DFTVKIGDFGMTRDVYETDYYRKGGKGLLP VRWMAPESLKDGIFTTHSDVWSFGVLWEIV TLAEQPYQGLSNEQVLKFVMDGGVLEELEGC PLQLQELMSRCWQPNPRLRPSFTHILDSIQEEL RPSFRLLSFYYSPECRGARGSLPTTDAEPDSSP
723	2073	A	5672	1	216	TPRDCSPQNGGPGH LÄWIDNILPEKEKKETDKKRKRKKGAHEDCD EEPQFPPPSVIKIPMESVQSDPQNGIHCIARKR SSSWSYSL
724	2074	A	5704	4235	940	ARGRRSRPVWAASWGGRGRPAARRRPRGLA ATMGFELDRFDGDVDPDLKCALCHKVLEDP LTTPCGHVFCAGCVLPWVVQEGSCPARCRGR LSAKELNHVLPLKRLILKLDIKCAYATRGCGR VVKLQQLPEHLERCDFAPARCRHAGCGQVLL RRDVEAHMRDACDARPVGRCQEGCGLPLTH GEQRAGGHCCARALRAHNGALQARLGALHK ALKEALRAGKREKSLVAQLAAAQLELQMT ALRYQKKFTEYSARLDSLSRCVAAPPGGKGE ETKSLTLVLHRDSGSLGFNIIGGRPSVDNHDG SSSEGIFVSKIVDSGPAAKEGGLQIHDRIIEVN GRDLSRATHDQAVEAFKTAKEPIVVQVLRRT PRTKMFTPPSESQLVDTGTQTDITFEHIMALT KMSSPSPPVLDPYLLPEEHPSAHEYYDPNDYI GDIHQEMDREELELEEVDLYRMNSQDKLGLT VCYRTDDEDDIGIYISEIDPNSIAAKDGRIREG DRIIQINGIEVQNREEAVALLTSEENKNFSLLI ARAELQI.DEGWMDDDRNDFLDDLHMDMLE EQHHQAMQFTASVLQQKKHDEDGGTTDTAT ILSNQHEKDSGVGRTDESTRNDESSEQENNG DDATASSNPLAGQRKLTCSQDTLGSGDLPFS NESFISADCTDADYLGIPVDECERFRELLELK CQVKSATPYGLYYPSGPLDAGKSDPESVDKE LELLNEELRSIELECLSIVRAHKMQQLKEQYR ESWMLHNSGFRNYNTSIDVRRHELSDITELPE KSDKDSSSAYNTGESCRSTPLTLEISPDNSLRR

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						SRSPTPSQKLGSAYLPSYHHSPYKHAHIPAHA QHYQSYMQLIQQKSAVEYAQSQMSLVSMCK DLSSPTPSEPRMEWKVKIRSDGTRYITKRPVR DRLLRERALKIREERSGMTTDDDAVSEMKM GRYWSKEERKQHLVKAKEQRRRREFMMQSR LDCLKEQQAADDRKEMNILELSHKKMMKKR NKKIFDNWMTIQELLTHGTKSPDGTRVYNSF LSVTTV
725	2075	A	5707	3	1770	QISTEVSEAPVANDKPKTLVVKVQKKAADLP DRDTWKGRFDFLMSCVGYAIGLGNVWRFPY LCGKNGGGAFLIPYFLTLIFAGVPLFLLECSLG QYTSIGGLGVWKLAPMFKGVGLAAAVLSFW LNIYYIVIISWAIYYLYNSFTTTLPWKQCDNP WNTDRCFSNYSMVNTTNMTSAVVEFWERN MHQMTDGLDKPGQIR WPLAITLAIAWILVYF CIWKGVGWTGKVVYFSATYPYIMLIILFFRGV TLPGAKEGILFYITPNFRKLSDSEVWLDAATQ IFFSYGLGLGSLIALGSYNSFHNNVYRDSIIVC CINSCTSMFAGFVIFSIVGFMAHVTKRSIADV AASGPGLAFLAYPEAVTQLPISPLWAILFFSM LLMLGIDSQFCTVEGFITALVDEYPRLLRNRR ELFIAAVCIISYLIGLSNITQGGIYVFKLFDYYS ASGMSLLFLVFFECVSISWFYGVNRFYDNIQE MVGSRPCIWWKLCWSFFTPIIVAGVFIFSAVQ MTPLTMGNYVFPKWGQGVGWLMALSSMVL IPGYMAYMFLTLKGSLKQRIQVMVQPSEDIV RPENGPEQPQAGSSTSKEAYI
726	2076	A	5711	156	423	PRRDPGRTPELRGSAPRKTGANMPVRRGHVA PQNTFLGTIIRKFEGQNKKFIIANARVQNCAII YCNDGFCEMTGFSRPDVMQKPCTCD
727	2077	A	5716	3	274	HASEYFFKLCSFQVFLSFPLATIVIDVGLVVIP LVKSPNVHYVYVLLLVLSGLLFYIPLIHFKIRL AWFEKMTCYLQLLFNICLPDVSEE
728	2078	A	5737	1899		IQASRASPYPRVKVDFALSCHEDLLAPISEPIE WKYHSPEEEISLGPACWLWDFLRRSQQAGFL LPLSGGVDSAATACLIYSMCCQVCEAVRSGN EEVLADVRTIVNQISYTPQDPRDLCGRILTTC YMASKNSSQETCTRARELAQQIGSHHISLNID PAVKAVMGIFSLVTGKSPLFAAHGGSSRENL ALQNVQARIRMVLAYLFAQLSLWSRGVHGG LLVLGSANVDESLLGYLTKYDCSSADINPIGG ISKTDLRAFVQFCIQRFQLPALQSILLAPATAE LEPLADGQVSQTDEEDMGMTYAELSVYGKL RKVAKMGPYSMFCKLLGMWRHICTPRQVAD KVKRFFSKYSMNRHKMTTLTPAYHAENYSPE DNRFDLRPFLYNTSWPWQFRCIENQVLQLER AEPQSLDGVD
729	2079	A	5741	1	5976	PGCAARLSRARAPGPGAAGAGRKRLADPGPP PASRRLRAPGSRPRLAPCTRRAAQPAHARMA PRAAGGÁPLSARAAAASPPPFQTPPRCPVPLL LLLLLGAARAGALEIQRRFPSPTPTNNFALDG AAGTVYLAAVNRLYQLSGANLSLEAEAAVG PVPDSPLCHAPQLPQASCEHPRRLTDNYNKIL QLDPGQGLVVVCGSIYQGFCQLRRRGNISAV AVRFPPAAPPAEPVTVFPSMLNVAANHPNAS TVGLVLPPAAGAGGSRLLVGATYTGYGSSFF PRNRSLEDHRFENTPEIAIRSLDTRGDLAKLFT

730 2080 A 5744 3 292 QPSPLFHSHLETLQLLRTAQLPEQVSWPWGQ VANGKGNQRNMGSPQPSLLAFERNLELQIMG LGYSLLMGKLRPRVAKDTLRVHRDSTPSPLT LKD	SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion FDLNPSDDNILKIKQGAKEQHKLGFVSAFLHP SDPPPGAQSYAYLALNSEARAGDKESQARSL LARICLPHGAGGDAKKLTESYIQLGLQCAGG AGRGDLYSRLVSVFPARERLFAVFERPQGSPA ARAAPAALCAFRFADVRAAIRAARTACFVEP APDVVAVLDSVVQGTGPACERKLNIQLQPEQ LDCGAAHLQHPLSILQPLKATPVFRAFGLTSV AVASVNNYTAVFLGTVNGRLLKINLMESMQ VVSRRVVTVAYGEPVHHVMQFDPADSGYLY LMTSHQMARVKVAACNVHSTCGDCVGAAD AYCGWCALETRCTLQQDCTNSSQQHFWTSA SEGPSRCPAMTVLPSEIDVRQEYPGMILQISGS LPSLSGMEMACDYGNNIRTVARVPGPAFGHQ IAYCNLLPRADGFPPFPPNQDHVTYEMSVRVN GRNIVKANFTIVDCSRTAQVYPHTACTSCLSA QWPCFWCSQQHSCVSNQSRCEASPNPTSPQD CPRTLLSPLAPVTGGSQNILVPLANTAFFQG AALECSFGLEEIFEAVWNESVVRCDQVUH TTRKSQVFPLSQLKGRRAFFLDSPEPMTVM VYNCAMGSPDCSQCLGREDLGHLCMWSDGC RLRGPLQPMAGTCPAPEIRAIEPLSGPLDGGT LLTITIGRRILGRRLSDVAHGWIGGVACEPLP DRYTVSEEIVCVTGPAPGPLSGVTVNASKE GKSRDRFSYVLPLVHSLEPTMGPKAGGTRITI HGNDLHVGSELQVLVNDTDPCTELMRTDTSI ACTMPEGALPAPVPCVCVRFERRGCVHGNLTF WYMQNPVITAISPRSPVSGGRTITVAGERFH MVQNVSMAVHHIGREPTLCKVLNSTLITCPSP GALSNASAPVDFFINGRAYADEVAVELLUNITLITCPSP GALSNASAPVDFFINGRAYADEVAVELLUNITLITCPSP GALSNASAPVDFFINGRAYADEVAVELLUNITLITCPSP GALSNASAPVDFFINGRAYADEVAVELLUNITLITCPSP GALSNASAPVDFFINGRAYADEVAVELLUNITLITCPSP GALSNASAPVDFFINGRAYADEVAVELLUNITLITCPSP GALSNASAPVDFFINGRAYADEVAVELLUNITLITCPSP GALSNASAPVDFFINGRAYADEVAVELLUNITLARGE GISLFSSLLDNKHELIVFVHALEQQKDFAVRD RCSLASLLTIALHGKLEYTSIMKELLVDLID ASAAKNPKLMLRTESVVEKMLTNWMSICM YSCLRETVGEPFFLLLCAIKQQINKGSIDAITG KARYTLNEEWLLRENIEAKFRANUNSCQCGGMDSLSVARAMDTDTLTQVREKILEAFCKNVPY SQWPRAEDVDLEWFASSTQSYILRDLDDTSV VEDGRKLNTLAHYKLPEGASLAMSLIDKKD NTLGRVKDLDTEKYFHLLVPTDELAEPKKSH RQSHRKKVLPETYTRLLSTKGTLQKFLDDTF KAILSIREDKPPLAVKYFFDFLEEQAEKRGISD PDTLHIWKTNSLPLRFWVNILKNPQFVFDIDK TDHILGRVKDLDTEKYFHUNDAALEKSPRANTADETOLAUKARDENAMAEIYXYAK PUDBOALBESRKYQNETNTNV
730 2080 A 5744 3 292 QPSPLFHSHLETLQLLRTAQLPEQVSWPWGQ VANGKGNQRNMGSPQPSLLAFERNLELQIMG LGYSLLMGKLRPRVAKDTLRVHRDSTPSPLT LKD							RYRPQIMAALEANPTARRTQLQHKFEQVVAL
LGYSLLMGKLRPRVAKDTLRVHRDSTPSPLT LKD	730	2080	A	5744	3	292	QPSPLFHSHLETLQLLRTAQLPEQVSWPWGQ
							LGYSLLMGKLRPRVAKDTLRVHRDSTPSPLT
731   2081   A   5747   1   382   FLKCMRKAFRSSKLLQVGYTPDGKDDYRWC   FRVDEVNWTTWNTNVGIINEDPGNCEGVKRT	731	2081	A	5747	1	382	FLKCMRKAFRSSKLLQVGYTPDGKDDYRWC

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732	2082	A	5753	198	3	RQSAQPEEVYLRQFSGSLKPEDAEVFKSPAAS GEK AQAESSTVASPEATAGPLCTRIPNVPPPTPIRP
733	2083	A .	5754	2	2223	PGKLQAQLPCPSPVRFTSARIPPASRPQTKS  AAGPPGLEAEGRAPESAGPGPGGDAAETPGL PPAHSGTLMMAFRDVTVQIANQNISVSSSTAL SVANCLGAQTVQAPAEPAAGKAEQGETSGR EAPEAPAVGREDASAEDSCAEAGASGAADG ATAPKTEEEEEEEETAEVGRGAEAEAGDLEQ LNRTSTSTKSAKSGSEASASASKDALQAMILS LPRYHCENPASCKSPTLSTDTLRKRLYRIGLN LFNINPDKGIQFLISRGFIPDTPIGVAHFLLQRK
						GLSRQMIGEFLGNSKKQFNRDVLDCVVDEM DFSSMELDEALRKFQAHIRVQGEAQKVERLIE AFSQRYCMCNPEVVQQFHNPDTIFILAFAIILL NTDMYSPNIKPDRKMMLEDFIRNLRGVDDG ADIPRELVVGIYERIQQKELKSNEDHVTYVTK VEKSIVGMKTVLSVPHRRLVCCSRLFEVTDV NKLQKQAAHQREVFLFNDLLVILKLCPKKKS SSTYTFCKSVGLLGMQFQLFENEYYSHGITLV TPLSGSEKKQVLHFCALGSDEMQKFVEDLKE SIAEVTELEQIRIEWELEKQQGTKTLSFKPCGA QGDPQSKQGSPTAKREAALRERPAESTVEVSI HNRLQTSQHNSGLGAERGAPVPPPDLQPSPPR QQTPPLPPPPPTPPGTLVQCQQIVKVIVLDKPC LARMEPLLSQALSCYTSSSSDSCGSTPLGGPG SPVKVTHQPPLPPPPPPYNHPHQFCPPGSLLH GHRYSSGSRSLV
734	2084	A	5788	8	362	SSVMGDLVGQGLEEQIVARDENSWLIDGGTP IDDVMRVLDIDEFPQSGNYETIGGFMMFMLR KIPKRTDSVKFAGYKFEVVDIDNYRIDQLLVT RIDSKATALSPKLPDAKDKEESVA
735	2085	A	5827	1	1257	MVFSAVLTAFHTGTSNTTFVVYENTYMNITL PPPFQHPDLSPLLRYSFETMAPTGLSSLTVNST AVPTTPAAFKSLNLPLQITLSAIMIFILFVSFLG NLVVCLMVYQKAAMRSAINILLASLAFADM LLAVLNMPFALVTILTTRWIFGKFFCRVSAMF FWLFVIEGVAILLIISIDRFLIIVQRQDKLNPYR AKVLIAVSWATSFCVAFPLAVGNPDLQIPSRA PQCVFGYTTNPGYQAYVILISLISFFIPFLVILY SFMGILNTLRHNALRIHSYPEGICLSQASKLGL MGLQRPFQMSIDMGFKTRAFTTILILFAVFIVC WAPFTTYSLVATFSKHFYYQHNFFEISTWLL WLCYLKSALNPLIYYWRIKKFHDACLDMMP KSFKFLPQLPGHTKRRIRPSAVYVCGEHRTVV
736	2086	A	5870	3	268	FTRSDELARHYRTHTGEKRFSCPLCPKQFSRS DHLTKHARRHPTYHPDMIEYRGRRRTPRIDPP LTSEVESSASGSGPGPAPSFTTCL
737	2087	A	5871	2	521	LTWPQLFLETLPELLHMSRPAEDGPSPGALVR RSSSLGYISKAEEYFLLKSRSDLMFEKQSERH GLARRLTTARRPPASSEQAQQELFNELKPAV DGANFIVNHMRDQNNYNEEKDSWNRVART VDRLCLFVVTPVMVVGTAWIFLQGVYNQPPP QPFPGDPYSYNVQDKRFI
738	2088	A	5881	1	1160	LVVTAITAILAFPNEYTRMSTSELISELFNDCG LLDSSKLCDYENRFNTSKGGELPDRPAGVGV YSAMWQLALTLILKIVITIFTFGMKIPSGLFIPS MAVGAIAGRLLGVGMEQLAYYHQEWTVFNS

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739	2089	A	5892	2	916	LVTHNGRLLGIITKKDVLKHIAQMANQDPDSI LFN TLQLAASVPFFAISLISWWLPESARWLIINGKP DQALQELRKVARINGHKEAKNLTIEVLMSSV KEEVASAKEPRSVLDLFCVPVLRWRSCAMLV VNFSLLISYYGLVFDLQSLGRDIFLLQALFGA
						VDFLGRATTALLLSFLGRRTIQAGSQAMAGL AILANMLVPQDLQTLRVVFAVLGKGCFGISL TCLTIYKAELFPTPVRMTADGILHTVGRLGA MMGPLILMSRQALPLLPPLLYGVISIASSLVVL FFLPETQGLPLPDTIQDLESQKSTAAQGNRQE AFTVESTSLLEIVALHGAL
740	2090	A	5900	2	426	RPIKTLGIGFHFSVDGVHFLTQREVQNLWKE NLIILDTAKKHGYEVVDTFTITMGRYKEFLQG KCGCHFHEVVKSKLSKEYNFIKMKRSRNHIM GRYFSNQSKLQQGTVTNFRSPYHVRGPINQV CSEILLSRMCANKRTM
741	2091	A	5910	3	412	RMPESTLLIICENGYILEAPLPTIKQEEDDHDV VSYEIKDMCIKCFHFSSVKSKILRLIEIEKRER QRELKEKIREERRNKLAAEMGEDGEKEFQEE EEEKEEEEEEEPLPEIFIPSTPSPILCGFYSEPG KFWV
742	2092	A	5936	1	482	MGCRLLCCVVFCLLQAGPLDTAVSQTPKYLV TQMGNDKSIKCEQNLGHDTMYWYKQDSKK FLKIMFSYNNKELIINETVPNRFSPKSPDKAHL NLHINSLELGDSAVYFCASSQDTALQSHCIPV HKPPGSARKLQGSVCTCTQGSSLHSLMASDG VPVC
743	2093	A	5938	Ī	1566	MNSFFGTPAASWCLLESDVSSAPDKEAGRER RALSVQQRGGPAWSGSLEWSRQSAGDRRRL GLSRQTAKSSWSRSRDRTCCCRRAWWILVPA ADRARRERFIMNEKWDTNSSENWHPIWNVN DTKHHLYSDINITYVNYYLHQPQVAAIFIISYF LIFFLCMMGNTVVCFIVMRNKHMHTVTNLFI LNLAISDLLVGIFCMPI'ILLDNIIAGWPFGNTM CKISGLVQGISVAASVFTLVAIAVDRFQCVVY PFKPKLTIKTAFVIIMIIWVLAITIMSPSAVMLH VQEEKYYRVRLNSQNKTSPVYWCREDWPNQ EMRKIYTTVLFANIYLAPLSLIVIMYGRIGISLF RAAVPHTGRKNQEQWHVVSRKKQKIIKMLLI VALLFILSWLPLWTLMMLSDYADLSPNELQII NIYIYPFAHWLAFGNSSVNPIIYGFFNENFRRG FQEAFQLQLCQKRAKPMEAYALKAKSHVLIN TSNQLVQESTFQNPHGETLLYRKSAEKPQQE LVMEELKETTNSSEI
744	2094	A	5966	149	327	SHVCVSHYAGSSGCPAGAGAGAVALGISAVA LYDYQGGRLGVARGAWYMEAPDIRQGDM
745	2095	A	5970	413	856	GAPHTDWAWAPTPMSGLGSGRGRQGTLASS PLSLPLLLAGVTGILATELFDQMARPAACMV CGALMWIMLILVGLGFPFIMEALSHFLYVPFL GVCVCGAIYTGLFLPETKGKTFQEISKELHRL NFPRRAQGPTWRSLEVIQSTEL

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746	2096	A	5971	3	1343	AQTARRIIGLELDTEGHRLFVAFSGCIVYLPLS RCARHGACQRSCLASQDPYCGWHSSRGCVDI RGSGGTDVDQAGNQESMEHGDCQDGATGSQ SGPGDSAYGVRRDLPPASASRSVPIPLLLASV AAAFALGASVSGLLVSCACRRAHRRRGKDIE TPGLPRPLSLRSLARLHGGGPEPPPSKDGDA VQTPQLYTTFLPPPEGVPPPELACLPTPESTPE LPVKHLRAAGDPWEWNQNRNNAKEGPGRSR GGHAAGGPAPRVLVRPPPPGCPGQAVEVTTL EELLRYLHGPQPPRKGAEPPAPLTSRALPPEP APALLGGPSPRPHECASPLRLDVPPEGRCASA PARPALSAPAPRLGVGGGRRLPFSGHRAPPAL LTRVPSGGPSRYSGGPGKHLLYLGRPEGYRG RALKRVDVEKPQLSLKPPLVGPSSRQAVPNG GRFNF
747	2097	A	5998	2	754	DHASLPCSWNHRFDVETRHVFIGDHSGQVTI LKLEQENCTLVTTFRGHTGGVTALCWDPVQ RVLFSGSSDHSVIMWDIGGRKGTAIELQGHN DRVQALSYAQHTRQLISCGGDGGIVVWNMD VERQETPEWLDSDSCQKCDQPFFWNFKQMW DSKKIGLRQHHCRKCGKAVCGKCSSKRSSIPL MGFEFEVRVCDSCHEAITDEERAPTATFHDSK HNIVHVHFDATRGWLLTSGTDKVIKLWDMT PVVS
748	2098	Α	6001	2	747	AMVFGGVVPYVPQYRDIRRTQNADGFSTYV CLVLLVANILRILFWFGRRFESPLLWQSAIMIL TMLLMLKLCTEVRVANELNARRRSFTAADS KDEEVKVAPRRSFLDFDPHHFWQWSSFSDYV QCVLAFTGVAGYITYLSIDSALFVETLGFLAV LTEAMLGVPQLYRNHRHQSTEGMSIKMVLM WTSGDAFKTAYFLLKGAPLQFSVCGLLQVLV DLAILGQAYAFARHPQKPAPHAVHPTGTKAL
749	2099	A	6002	2	447	GRPDRSELVRMHILEETFAEPSLQATQMKLK RARLADDLNEKIAQRPGPMELVEKNILPVDSS VKEAIIGVGKEDYPHTQGDFSFDEDSSDALSP DQPASQESQGSAASPSEPKVSESPSPVTTNTP AQFASVSPTVPEFLKTPPTAD
750	2100	A	6004	2	427	LLTQAMLVLPHRPQWFTPGPRLQAQGPCQEG WRWELRLRNYVPEDEDLNKRRVPQAKPDAV QEKVKEQLEAAKPEPVIEEVDLAKLAPRKPD WDLKRDVAKKLEKLLKRTQRAIAELIRERLK GQEDSLDSAVDAATEHKTC
751	2101	A	6007	,	1280	TDQAKVDNQPEKLVRSAEDVSTVPTQPDNPF SHPDKLKRMSKSVPAFLQDESDDRETDTASE SSYQLSRHKKSPSSLTNLSSSSGMTSLSSVSGS VMSVYSGDFGNLEVKGNIQFAIEYVESLKEL HVFVAQCKDLAAADVKKQRSDPYVKAYLLP DKGKMGKKKTLVVKKTLNPVYNEILRYKIEK QILKTQKLNLSIWHRDTFKRNSFLGEVELDLE TWDWDNKQNKQLRWYPLKRKTAPVALEAE NRGEMKLALQYVPEPVPGKKLPTTGEVHIWV KECLDLPLLRGSHLNSFVKCTILPDTSRKSRQ KTRAVGKTTNPIFNHTMVYDGFRPEDLMEAC VELTVWDHYKLTNQFLGGLRIGFGTGKSYGT EVDWMDSTSEEVALWEKMVNSPNTWIEATL PLRMLLIAKISK
752	2102	A	6028	108	1283	KEIFSPFELISVKPLCLLLGVTCSQSMAFEELL SQVGGLGRFQMLHLVFILPSLMLLIPHILLENF AAAIPGHRCWVHMLDNNTGSGNETGILSEDA

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753	2103	A	6043	1	1470	DSFESILRLIFEIHHSGEKGDIVVFLACEQDIEK VCETVYQGSNLNPDLGELVVVPLYPKEKCSL FKPLDETEKRCQVYQRRVVLTTSSGEFLIWSN SVRFVIDVGVERRKVYNPRIRANSLVMQPISQ SQAEIRKQILGSSSSGKFFCLYTEEFASKDMTP LKPAEMQEANLTSMVLFMKRIDIAGLGHCDF MNRPAPESLMQALEDLDYLAALDNDGNLSE FGIIMSEFPLDPQLSKSILASCEFDCVDEVLTIA AMVTAPNCFSHVPHGAEEAALTCWKTFLHPE GDHFTLISIYKAYQDTTLNSSSEYCVEKWCRD YFLNCSALRMADVIRAELLEIIKRIELPYAEPA FGSKENTLNIKKALLSGYFMQIARDVDGSGN YLMLTHKQVAQLHPLSGYSITKKMPEWVLF HKFSISENNYIRITSEISPELFMQLVPQYYFSNL PPSESKDILQQVVDHLSPVSTMNKEQQMCET CPETEQRCTLQ
754	2104	A	6055	2	394	YYALHHWPFPDLLCQTTGAIFQMNMYGSCIF LMLINVDRYAAIVHPLRLRHLRRPRVARLLC LGVWALILVFAVPAARVHRPSRCRYRDLEVR LCFESFSDELWKGRLLPLVLLAEALGFLLPLA AVVYSS
755	2105	A	6059		1795	LGLGSGTLLSVSEYKKKYREHVLQLHARVKE RNARSVKITKRFTKLLIAPESAAPEEALGPAEE PEPGRARRSDTHTFNRLFRRDEEGRRPLTVVL QGPAGIGKTMAAKKILYDWAAGKLYQGQVD FAFFMPCGELLERPGTRSLADLILDQCPDRGA PVPQMLAQPQRLLFILDGADELPALGGPEAAP CTDPFEAASGARVLGGLLSKALLPTALLLVTT RAAAPGRLQGRLCSPQCAEVRGFSDKDKKK YFYKFFRDERRAERAYRFVKENETLFALCFV PFVCWIVCTVLRQQLELGRDLSRTSKTTTSVY LLFITSVLSSAPVADGPRLQGDLRNLCRLARE GVLGRRAQFAEKELEQLELRGSKVQTLFLSK KELPGVLETEVTYQFIDQSFQEFLAALSYLLE DGGVPRTAAGGVGTLLRGDAQPHSHLVLTT RFLFGLLSAERMRDIERHFGCMVSERVKQEA LRWVQGQGQCPGVAPEVTEGAKGLEDTEE PEEEEEGEEPNYPLELLYCLYETQEDAFVRQA LCRFPELALQRVRFCRMDVAVLSYCVRCCPA GQALRLISCRLVAAQEKKKKSLGKRLQASLG GG
756	2106	<b>A</b>	6060	12	436	SGRPTRPAKPTGQGMGRFMLTLVCQGSIMMS ARDLIMNNLTELQPGLFHHLRFLEELRLSGNH LSHIPGQAFSGLYSLKILMLHNNQLGGIPAQA LWELPSLQSLRLDANLISLVPERSFEGLSSLRH LWLDDNALTEIPS
757	2107	A	6063	54	419 .	ITPLGLGAADMCAFPWLLLLLLLQEGSQRRL WRWCGSEEVVAVLQESISLPLEIPPDEEVENII WSSHKSLATVVPGKEGHPATIMVTNPHYQG

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758	2108	A	6066	125	438	IGISCPATIFVPMFSHSLIGIGEEYQLPYYNMV PSDPSYEDMREVVCVKRLRPIVSNRWNSDEC LRAVLKLMSECWAHNPASRLTALRIKKTLAK MVESQDVK1
759	2109	A	6072	3	650	PGRRFRPAALEERAMEKLREKVPFQNRGKGT LSSIIPNNSDTRKATETTSLSSKPEYVNPDFRW SKDPSSKSGNLLETSEVGWTSNPEELDPIRLA LLGKSGLSCQVGSATSHPVSCQEPIDEDQRISP KDKSTAGREFSGQVSHQTTSENQCTPIPSSTV HSSVADMQNMPAAVHALLTQPSLSAAPFAQ RYLGTLPSTGSTTLPQCHAGNATVW
760	2110	A	6077	3	730	PLRLTLMEEVILLGLKDREGYTSFWNDCISSG LRGCMLIELPLRGRLQLEACGMRRKSLLTRK VICKSDAPTGDVLLDEALKHVKETQPPETVQ NWIELLSGETWNPLKLHYQLRNVRERLAKNL VEKGVLTTEKQNFLLFDMTTHPLTNNNIKQR LIKKVQEAVLDKWVNDPHRMDRRLLALIYL AHASDVLENAFAPLLDEQYDLATKRVRQLLD LDPEVECLKANTNEVLWAVVAAFTK
761	2111	A	6078	833	390	IVSFHLSGFKKFVRPFSFLSVHGLQVDEYHSV HQKLSADMADHSNLIRSLLVGAEDARLMRD MKTMKSRYMELYDLNRDLLNGYKIRWNNH TELLGNLKAVNQAIQRAGRLRVGKPKNQVIT ACRDAIRSNNINTLFKIMRVGTASS
762	2112	A	6079	2	2686	KKAITCGEKEKQDLIKSLAMLKDGFRTDRGS HSDLWSSSSSLESSSFPLPKQYLDVSSQTDISG SFGINSNNQLAEKVRLRLRYEEAKRIANLKI QLAKLDSEAWPGVLDSERDRLILINEKEELLK EMRFISPRKWTQGEVEQLEMARKRLEKDLQ AARDTQSKALTERLKLNSKRNQLVRELEEAT RQVATLHSQLKSLSSSMQSLSSGSSPGSLTSSR GSLVASSLDSSTSASFTDLYYDPFEQLDSELQ SKVEFLLLEGATGFRPSGCITTIHEDEVAKTQ KAEGGGRLQALRSLSGTPKSMTSLSPRSSLSS PSPPCSPLMADPLLAGDAFLNSLEFEDPELSA TLCELSLGNSAQERYRLEEPGTEGKQLGQAV NTAQGCGLKVACVSAAVSDESVAGDSGVYE ASVQRLGASEAAAFDSDESEAVGATRIQIALK YDEKNKQFAILIIQLSNLSALLQQDQKVNIR VAVLPCSESTTCLFRTRPLDASDTLVFNEVFW VSMSYPALHQKTLRVDVCTTDRSHLEECLGG AQISLAEVCRSGERSTRWYNLLSYKYLKKQS RELKPVGVMAPASGPASTDAVSALLEQTAVE LEKRQEGRSSTQTLEDSWRYEETSENEAVAE EEEEEVEEEGEEDVFTEKASPDMDGYPALK VDKETNTETPAPSPTVVRPKDRRVGTPSQGPF LRGSTIIRSKTFSPGPQSQYVCRLNRSDSDSST LSKKPPFVRNSLERRSVRMKRPSPPPQPSSVK SLRSERLIRTSLDLELDLQATRTWHSQLTQEIS VLKELKEQLEQAKSHGEKELPQWLREDERFR LLLRMLEKRMDRAEHMGELQTDKMMRAAA KDVHRLRGQSCKEPPEVQSFREKMAFFTRPR
763	2113	A	6082	3	1558	PHPIRFSKLCVSFNNQEYNQFCVIEEASKANE VLENLTQGKMCLVPGKTRKLLFKFVAKTED VGKKIEITSVDLALGNETGRCVVLNWQGGGG DAASSQEALQAARSFKRRPKLPDNEVHWGSII IQASTMIISRVPNISVHLLHEPPALTNEMYCLV

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764	2114	A	6093	1	1422	DDTSIAAA  AAADLANSNAGAAVGRKAGPRSPPSAPAPAP PPPAPAPPTLGNNHQESPGWRCCRPTLRERN ALMFNNELMADVHFVVGPPGATRTVPAHKY VLAVGSSVFYAMFYGDLAEVKSEIHIPDVEPA AFLILLKYMYSDEIDLEADTVLATLYAAKKYI VPALAKACVNFLETSLEAKNACVLLSQSRLF EEPELTQRCWEVIDAQAEMALRSEGFCEIDR QTLEIIVTREALNTKEAVVFEAVLNWAEAEC KRQGLPITPRNKRHVLGRALYLVRIPTMTLEE FANGAAQSDILTLEETHSIFLWYTATNKPRLD FPLTKRKGLAPQRCHRFQSSAYRSNQWRYRG RCDSIQFAVDRRVFIAGLGLYGSSSGKAEYSV KIELKRLGVVLAQNLTKFMSDGSSNTFPVWF EHPVQVEQDTFYTASAVLDGSELSYFGQEGM TEVQCGKVAFQFQCSSDSTNGTGVQGGQIPE LIFYA
765	2115	Α .	6099	1		SGFTHYAIYDFIVKGSCFCNVHADQCIPVHGF RPVKAPGTFHMVHGKCMCKHNTAGSHCQH CAPLYNDRPWEAADGKTGAPNECRTCKCNG HADTCHFDVNVWEASGNRSGGVCDDCQHN TEGQYCQRCKPGFYRDLRRPFSAPDACKPCS CHPVGSAVLPANSVTFCDPSNGDCPCKPGVA GRRCDRCMVGYWGFGDYGCRPCDCAGSCD PITGDCISSHTDIDWYHEVDFRPVHNKSEPP WEWEDAQGFSALLHSGKCECKEQTLGNAKA FCGMKYSYVLKIKILSAHDKGTHVEVNVKIK KVLKSTKLKIFRGKRTLYPESWTDRGCTCPIL NPGLEYLVAGHEDIRTGKLIVNMKSFVQHWK PSLGRKVMDILKRECK
766	2116	A	6103	2	384	MTAAATATVLKEGVLEKRSGGLLQLWKRKR CVLTERGLQLFEAKGTGGRPKELSFARIKAVE CVESTGRHIYFTLVTEGGGEIDFRCPLEDPGW NAQITLGLVKFKNQQAIQTVRARQSLGTGTL VS
767	2117	A	6106	1	542	SGSSHASDGSGFQELRICSEDQTPLIAGMCSLP MARYYIIKYADQKALYTRDGQLLVGDPVAD NCCAEKICTLPNRGLDRTKVPIFLGIQGGSRC LACVETEEGPSLQLEDVNIEELYKGGEEATRF TFFQSSSGSAFRLEAAAWPGWFLCGPAEPQQ PVQLTKESEPSARTKFYFEQSW
768	2118	A	6109	3	292	FILQAVLQLSSQEARYKAFGTCVSHIGAILAF YTPSVISSVMHRVARCAAPHVHILLANFYLLF PPMVNPIIYGVKTKQIRDSLGSIPEKGCVNRE
769	2119	A	6110	1	711	RHEPSCSNGVASTKSKQNHSKYPAPSSSSSSS SSSSSSSSSSVNYSESNSTDSTKSQHHSSTSNQ ETSDSEMEMEAEHYPNGVLGSMSTRIVNGAY KHEDLQTDESSMDDRHPRRQLCGGNQAATE

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SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
						RIILFGRELQALSEQLGREYGKNLAHTEMLQD AFSLLAYSDPWSCPVGQQLDPIQREPVCAAL NSAILESQNLPKQPPLMLALGQASECLRLMA RAGLGSCSFARVDDYLH
770	2120	A	6125	2	570	YFGLNLHVQHLGNNVFLLQTLFGAVILLANC VAPWALKYMNRRASQMLLMFLLAICLLAIIF VPQEMQMLREVLATLGLGASALANTLAFAH GNEVIPTIIRARAMGINATFANIAGALAPLMM ILSVYSPPLPWIIYGVFPFISGFAFLLLPETRNK PLFDTIQDEKNERKDPREPKQEDPRVEVTQF
771	2121	A	6126	909	353	RSFVLDTASAICNYNAHYKNHPKYWCRGYF RDYCNIIAFSPNSTNHVALRDTGNQLIVTMSC LTKEDTGWYWCGIQRDFARDDMDFTELIVT DDKGTLANDFWSGKDLSGNKTRSCKAPKVV RKADRSRTSILIICILITGLGIISVISHLTKRRRS QRNRRVGNTLKPFSRVLTPKEMAPTEQM
772	2122	A	6148	7	810	FVLGILALSHTISPFMNKFFPASFPNRQYQLLF TQGSGENKEEIINYEFDTKDLVCLGLSSIVGV WYLLRKHWIANNLFGLAFSLNGVELLHLNN VSTGCILLGGLFIYDVFWVFGTNVMVTVAKS FEAPIKLVFPQDLLEKGLEANNFAMLGLGDV VIPGIFIALLLRFDISLKKNTHTYFYTSFAAYIF GLGLTIFIMHIFKHAQPALLYLVPACIGFPVLV ALAKGEVTEMFSYEESNPKDPAAVTESKEGT EASASKGLEKKEK
773	2123	A	6161	3	1088	CQPMLVTRKNHPKLLLRRTESVAEKMLTNW FTFLLYKFLKESAGEPLFMLYCAIKHQMEKG PIDAITGEARYSLSEDKLIRHLIDYKTLTLNCV NPENENAPEVPVKGLDCDTGTQAKEKLLDA AYKGVPYSQRPKAADMDLEWRQGRMARIIL QDEDVTTKIDNDWKRLNTLAHYQVTDGSSV ALVPKQTSAYNISNSSTFTKSLSRYESMLRTA SSPDSLRSRTPMITPDLESGTKLWHLVKNHDH LDQREGDRGSKMVSEIYLTRLLATKGTLQKF VDDLFETIFSTAHRGSALPLAIKYMFDFLDEQ ADKHQIHDADVRHTWKSNCLPLRFWVNVIK NPQFVFDIHKNSITDACLSVV
774	2124	A	6163		125	KTAVKKRNLNPVFNETLRYSVPQAELQGRVL SLSVWHRESLGRNIFLGEVEVPLDTWDWGSE PTWLPLQPRVPPSPDDLPSRGLLALSLKYVPA GSEGAGLPPSGELHFWVKEARDLLPLRAGSL DTYVQCFVLPDDSRASRQRTRVVRRSLSPVF NHTMVYDGFGPADLRQACAELSLWDHGALA NRQLGGTRLSLGTGSSYGLQVPWMDSTPEEK QLWQALLEQPCEWVDGLLPLRTNLAPRT
775	2125	A	6191	2	392	ARGIGSLGRDHSGSGGGTGMAGAWVRKAAD YVRSKDFRDYLMSTHFWGPVANWGLPIAAIT DMK\KSPEIISRRMTFAL*CYSLTFVRFAHYVQ \PWNWLMLGCHTAVDFDQLISSMPCISHGMT ASASAL
776	2126	A	6217	1	827	FRGYWGVREAFTDASWSGGLGPGKPGMKIT RQKHAKKHLGFFRNNFGVREPYQILLDGTFC QAALRGRIQLREQLPRYLMGETQLCTTRCVL KELETLGKDLYGAKLIAQKCQVRNCPHFKNA VSGSECLLSMVEEGNPHHYFVATQDQNLSVK VKKKPGVPLMFIIQNTMVLDKPSPKTIAFVKA VESG\RLSQCMRKKVSNISKRNRV**KTLNRG RRKKRKKISGPNPLSCLKKKKKAPDTQSSASE KKRKRKRIRNRSNPKVLSEKQNAEGE

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	L OFFICE TO	1-5 /	1-050		T	
SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence	J	}	914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
ĺ		1		amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
				peptide	sequence	/=possible nucleotide deletion, \=possible
			ļ	sequence		nucleotide insertion
777	2127	<del> </del>	6026		1402	
111	212/	A	6236	1038	1402	YYQISSLPSIVGNGIFLWLLICIFLAKQGGSRL*
Í		1		[	[	FQPFGRPRGGGHLRSGVLGQPGQHGETP/SFF
						YNSKISPALWGPPVIPSALGGEAGKSL*PRRQ
						RFQRGGIAPLPSRVRGRAKLFLKKK
778	2128	A	6237	422	913	ASFFHHHRGAFLLLLAIPGS*GQDQSLIHWSN
	ł	l				AVSNAD\LLDLK\N*LDH\LEEKMPL\EVKVVP
		1	l			PQVL\SEPN*RSGGCFSAPSFEVPPWTGEVKP/
						SPQRDGGALG\QGPLGIPSDSILALLKKQT*RA
						LLNWPLGSLRRSSCFGGQDGQDLKPRSGLGC
	ļ	<u> </u>				NSFRYRR
779	2129	A	6249	420	36	ARAPSPSFSVRDVELSDPARERGEMPVAVGP
İ		1				YGQSQPSCFDRVKMGFVMGCAVGMAAGAL
	ļ	1				FGTFSCLSSILVSSSG/SGMRGRELMGGIGKTM
						MQSGGTFGTFMAIGMGIRC*PWLPTTSVPSH
						OSOPMY
780	2130	A	6263	415	1380	RIMRMCDRGIQMLITTVGAFAAFSLMTIAVG
180	2130	ΙΛ.	0203	413	1360	
			l	1		TDYWLYSRGVCRTKSTSDNETSRKNEEVMT
		1				HSGLWRTCCLEGAFRGVCKKIDHFPEDADYE
						QDTAEYLLRAVRASSVFPILSVTLLFFGGLCV
1	ĺ	ĺ	i		ĺ	AASEFHRSRHNVILSAGIFFVSAGLSNIIGIIVYI
						S\ANAGRTPGQR\DSKKSYSYGWSF/YFSGAFS
		1	1			FIIGR/IIC*GVGLPWHIYIEKHQQLRAKSHSEF
			1	i		LKKSTFARLPPYRYRFRRRSSSRSTEPRSRDLS
		İ				PISKGFHTIPSTDISMFTLSRDPSKITMGTLLNS
			ì			DRDHAFLQFHNSTPKEFKESLHNNPANRRTT
1	1	ł	ł		}	PV
701	0101	<u> </u>	6074	600		
781	2131	A	6274	832	318	RIIKVKDLKQTLAIKTAYPRCKCLVEMDQIFH
		<b>l</b> .	į .			LQVKQKQLACLCTWQARDPDCPPSTKVVL/L
			1			VGPGMGCMVALFQDSIAWSNKSMPSSLSAIS
ĺ	[	1				QSPCQVQAPEGPSSFHLPTLSFTTCLSWQGGD
				ł		LEFLGDLKGCSELKNFQELITQSALVHPKADV
				1		WWYCGRPLLGTLPSN
782	2132	A	6281	1324	393	WISLPSSLLCRKNGSSAEDDRR\GEPSAEEAEG
102	2132	11	0261	1524	373	EREDWGIGSA*SVGAVSKVPSARF*RTYPS\E
	1	Í	ľ	ĺ		DEEEVTHQKSSSSDSNSEEHRKKKTSRSRNK
		1	1			KKRKNKSSKRKHRKYSDSDSNSESDTNSDSD
İ		1				DDKKRVKAKKKKKKKKKKKKKKKKKKKKKKKKKKKKKKKK
		1				ESSDSSCKDSEEDLSEATWMEQPNVADTMDL
1	Į.		1			IGPEAPIIHTSQDEKPLKYGHALLPGEGAAMA
	I	1				EYVKAGKRIPRRGEIGLTSEEIGSFECSGYVM
	I					SGSRHRRMEAVRLRKENQIYSADEKRALASF
1						NQEERRKRESKILASFREMVHKKTKGKDDK
783	2133	_	6305	201	1032	
103	2133	A	1 0303	201	1032	WDDYPQGALRREAAEGLHFLGPPGRVRGQ
	l	]	1			LRGITGPAWYCHSPSHSLLSAFCHLPTPSRCP
			1			AMARPPVPGSVVVPNWHES/RRGQGVPGLHS
						AQEPPAGVWAA*AASAAAA\LSIDTASYKIFV
	I					SGKSGVGKTALVAKLAGLEVPVVHHETTGIQ
	I	1				TTVVFWPAKLQASSRVVMFRFEFWDCGESA
1	1	1	1			LKKFDHMLLACMENTDAFLFLFSFTDRASFE
					'	DLPGQLARIAGEAPGVVRMVIGSKFDQYMHT
1	1	1	1			
	1010	<del> </del>			0.6	DVPERDLTAFRQAWELPLLRVKSVPGRRLG
784	2134	A	6308	86	96	GSSPDPASLITMKNQDKKNGAAKQSNPKSSP
1	ļ	}	}	l		GQPEAGPEGAQERPSQAAPAVEAEGPGSSQA
	1	1				PRKPEGAQARTAQSGALRDVSEELSRQLEDIL
	1					STYCVDNNQGGPGEDGAQGEPAEPEDAEKSR
	1					TYVARNGEPEPTPVVNGEKEPSKGDPNTEEIR
	1					QSDEVGDRDHRRPQEKKKAKGLGKEITLLM
		1	1			
]	1	J	J .			QTLNTLSTPEEKLAALCKKYAELLEEHRNSQ
		1	1			KQMKLLQKKQSQLVQEKDHLRGEHSKAVLA

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  RSKLESLCRELQRHNRSLKEEGVQRAREEEE KRKEVTSHFQVTLNDIQLQMEQHNERNSKLR QENMELAERLKKLIEQYELREEHIDKVFKHK DLQQQLVDAKLQQAQEMLKEAEERHQREKD FLLKEAVESQRMCELMKQQETHLKQOLALY
<b>70</b> 5						TEKFEEFQNTLSKSSEVFTTFKQEMEKMTKKI KKLEKETTMYRSRWESSNKALLEMAEEKTV RDKELEGLQVKIQRLEKLCRALQT/GAQ*PVR GQRWGSHRTSAVRIFS
785	2135	A	6319	1493	889	SPQGPLLRSVSPVSAGASSVTPGGAQPGVTTT PPSLVAVAPAPGSAAGPAAGWQ*HAGCR/WT KLPWSWGMRPMKIFFSEEYRSISTRISHDAL* EKCTQPAKPLSMIR\TGSSVSPG/PLVKWNWT RREFRNSGTRVVSSCCGMSCMYSFLGHCSV/S QDLPLVHVDVGWQPPLGPTVGLRPGLLPLHD TTPCQKLVVDDLDWA
786	2136	A	6320	551	135	RWLPVAECDSSCVGCTGEGPGNCKECISGYA REHGQCADVDECSLAEKTCVRKNENCYNTP GSYVCVCPDGFEET/RRCLCAAGRG*SHRRRK PDTAALPRRPVMCRTYPLNYSEGCPVENVAL RMPSPAVDSGGERLPAL
787	2137	A	6330	1693	227	DYVLTAELHRQRSPGVSFGLSVFNLMNAIMG SGILGLAYVMANTGVFGFSFLLLTVALLASYS VHLLLSMCIQTAYLGP*TNYFMVLPAH*LTCL PLIEFLQSL*NSL\*AVTSYEDLGLFAFGLPGKL VVAGTIIIQNIGAMSSYLLIIKTELPAAIAEFLT GDYSRYWYLDGQTLLIIICVGIVFPLALLPKIG FLGYTSSLSFFFMMFFALVVIIKKWSIPCPLTL NYVEKGFQISNVTDDCKPKLFHFSKESAYALP TMAFSFLCHTSILPIYCELQSPSKKRMQNVTN TAIALSFLIYFISALFGYLTFYD/GTTKAQRGE VTCHRIKDKVESELLKG***IP*SHDVVVMT\V KLCILFAVLL\TVPLIHFPARKAVTMMFFSNFP FSWIRHFLITLALNIIIVLLAIYVPDIRNVFGVV GASTSTCLIFIFPGLFYLKLSREDFLSWKKLGV GCFC/LLSFKTSILRNSLSVYIILPASRKSIYFKI
788	2138	A	6351	1	6622	PRSLCFSLWAEAAVLADGGLRRRRRLLRGTM SASFVPNGASLEDCHCNLFCLADLTGIKWKK YVWQGPTSAPILFPVTEEDPILSSFSRCLKADV LG/VWRRDQRPERRE\L*IFWGGEDP\VLLTLF TMTYQKKKMECGRMDFFMNAVLCFSKAVH NLLERCLMNRNFVRIGKWFVKPYEKDEKPIN KSEHLSCSFTFFLHGDSNVCTSVEINQHQPVY LLSEEHITLAQQSNSPFQVILCPFGLNGTLTGQ AFKMSDSATKKLIGEWKQFYPISCCLKEMSE EKQEDMDWEDDSLAAVEVLVAGVRMIYPAC FVLVPQSDIPTPSPVGSTHCSSSCLGVHQVPAS TRDPAMSSVTLTPPTSPEEVQTVDPQSVQKW VKFSSVSDGFNSDSTSHHGGKIPRKLANHVV DRVWQECNMNRAQNKRKYSASSGGLCEEAT AAKVASWDFVEATQRTNCSCLRHKNLKSRN AGQQGQAPSLGQQQQILPKHKTNEKQEKSEK PQKRPLTPFHHRVSVSDDVGMD\ADS\ASQRL V\ISAP\DSQ\VRFSNIR\TNDVAK\TPQMHGTE MANSPQPPPLSP\HPCDVVDEGVTKTPSTPQS QHFYQMPTPDFLVPSKPMEDRIDSLSQSFPPQ YQEAVEPTVYVGTAVNLEEDEANIAWKYYK FPKKKDVEFLPPQLPSDKFKDDPVGPFGQESV TSVTELMVQCKKPLKVSDELVQQYQIKNQCL SAIASDAEQEPKIDPYAFVEGDEEFLFPDKKD

SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-	]	USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence	ļ	09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence			914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan,
	Į			amino acid residue of	of peptide	Y=Tyrosine, X=Unknown, *=Stop codon,
	İ			peptide	sequence	/=possible nucleotide deletion, \=possible
	İ			sequence		nucleotide insertion
		<del> </del>	<u> </u>	sequence		
						RQNSEREAGKKHKVEDGTSSVTVLSHEEDA MSLFSPSIKQDAPRPTSHARPPSTSLIYDSDLA
						VSYTDLDNLFNSDEDELTPGSKRSANGSDDK
1	ĺ	ĺ	1			
						ASCKESKTGNLDPLSCISTADLHKMYPTPPSL EQHIMGFSPMNMNNKEYGSMDTTPGGTVLE
	1					GNSSSIGAQFKIEVDEGFCSPKPSEIKDFSYVY
						KPENCQILVGCSMFAPLKTLPSQYLPLIKLPEE
			1			CIYRQSWTVGKLELLSSGPSMPFIKEGDGSNM
						DQEYGTAYTPQTHTSCGMPPSSAPPSNSGAGI
						LPSPSTPRFPTPRTPRTPRTPRGAGGPASAQGS
1	}	ł	1	}		VKYENSDLYSPASTPSTCRPLNSVEPATVPSIP
						EAHSLYVNLILSESVMNLFKDCNSDSCCICVC
						NMNIKGADVGVYIPDPTQEAQYRCTCGFSAV
						MNRKFGNNSGLFFEDELDIIGRNTDCGKEAE
						KRFEALRATSAEHVNGGLKESEKLSDDLILLL
		1				QDQCTNLFSPFGAADQDPFPKSGVISNWVRV
		1				EERDCCNDCYLALEHGRQFMDNMSGGKVDE
	ļ	ļ	ļ			ALVKSSCLHPWSKRNDVSMQCSQDILRMLLS
						LQPVLQDAIQKKRTVRPWGVQGPLTWQQFH
						KMAGRGSYGTDESPEPLPIPTFLLGYDYDYLV
						LSPFALPYWERLMLEPYGSQRDIAYVVLCPE
		ļ				NEALLNGAKSFFRDLTAIYESCRLGOHRPVSR
			1			LLTDGIMRVGSTASKKLSEKLVAEWFSQAAD
						GNNEAFSKLKLYAQVCRYDLGPYLASLPLDS
		1				SLLSQPNLVAPTSQSLITPPQMTNTGNANTPS
1 1		(				ATLASAASSTMTVTSGVAISTSVATANSTLTT
		İ				ASTSSSSSSNLNSGVSSNKLPSFPPFGSMNSNA
						AGSMSTQANTVQSGQLGGQQTSALQTAGISG
	]					ESSSLPTQPHPDVSESTMDRDKVGIPTDGDSH
	E					AVTYPPAIVVYIIDPFTYENTDESTNSSSVWTL
						GLLRCFLEMVQTLPPHIKSTVSVQIIPCQYLLQ
						PVKHEDREIYPQHLKSLAFSAFTQCRRPLPTS
1 1	}	ł	1			TNVKTLTGFGPGLAMETALRSPDRPECIRLYA
						PPFILAPVKDKQTELGETFGEAGQKYNVLFV
						GYCLSHDQRWILASCTDLYGELLETCIINIDVP
	100	1	ļ			NRARRKKSSARKFGLQKLWEWCLGLVQMSS
		1				LPWRVVIGRLGRIGHGELKDWSCLLSRRNLQ
						SLSKRLKDMCRMCGISAADSPSILSACLVAM
						EPQGSFVIMPDSVSTGSVFGRSTTLNMQTSQL
] ]		ļ	ļ			NTPQDTSCTHILVFPTSASVQVASATYTTENL
						DLAFNPNNDGADGMGIFDLLDTGDDLDPDII
						NILPASPTGSPVHSPGSHYPHGGDAGKGQSTD
						RLLSTEPHEEVPNILQQPLALGYFVSTAKAGP
						LPDWFWSACPQAQYQCPLFLKASLHLHVPSV
		<b> </b>				QSDELLHSKHSHPLDSNQTSDVLRFVLEQYN
		}				ALSWLTCDPATQDRRSCLPIHFVVLNQLYNFI
700	0120		62.50	1	2002	MNML
789	2139	Α	6359	1	2002	TGTLTEDGLDVMGVVPLKGQAFLPLVPEPRR
		1				LPVGPLLRALATCHALSRLQDTPVGDPMDLK
			1			MVESTGWVLEEEPAADSAFGTQVLAVMRPP
						LWEPQLQAMEEPPVPVSVLHRFPFSSALQRM
1						SVVVAWPGATQPEAYVKGSPELVAGLCNPET
						VPTDFAQMLQSYTAAGYRVVALASKPLPSVP
[		1	[	[		SLEAAQQLTRDTVEGDLSLLGLLVMRNLLKP
						QTTPVIQALRRTRIRAVMVTGDNLQTAVTVA
	l	l				RGCGMVAPQEHLIIVHATHPERGQPASLEFLP MESPTAVNGVKDPDQAASYTVEPDPRSRHLA
						LSGPTFGIIVKHFPKLLPKVLVQGTVFARMAP
				-		

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  VIREGRCSLDTSFSVFKYMALYSLTQFISVLIL YTINTNLGDLQFLAIDLVITTTVAVLMSRTGP ALVLGRVRPPGALLSVPVLSSLLLQMVLVTG VQLGGYFLTLAQPWFVPLNRTVAAPDNLPNY ENTVVFSLSSFQYLILAAAVSKGAPFR\RPLTN NVPFLLASAL*SSVLVVLVLSPGLLHGPLALR NITDTGFKLLLVGLVTLNFVGGLHAGERARP VPPRLPAPPPAQAG\SKKRFKQLERELAEQPW
790	2140	A	6380	76	1059	PPLPAGPLR SSAGSARKLQVMALAARLWRLLPFRRGAAP GSRLPAGTSGSRGHCGPCRFRGFEVMGNPGT FKRGLLSALSYLGFETYQVISQAAVVHATA KVEEILEQADYLYESGETEKLYQLLTQYKESE DAELLWRLARASRDVAQLSRTSEEEKKLLVY EALEYAKRA/L/EKNESSFASHKWYAICLSDV GDYEGIKAKIANA YIIKEHFEKAIELNPKDATS IHLMGIWCYTFAEMPWYQRRIA*NACLQLPP *FPPYEKALG\YFHRAEQVDPNFYSKNLLLLG KTYLKLHNKKLAAFWLMKAKDYPAHTEED KQIQTEAAQLLTSFSEKN
791	2141		6434	3	1460	IALLIVDGLAWDDQGGLALLHISPSKLIL*QDS SGMS/YVMVRCTITRAFFKSLLCHICQYSIGPQ *VT\CPGQDACKE*KSTAN*GG*RE**PQVLFF AFLSNPAVKFGRMSKKQRDSLYAEVQKHQQ RLQEQRQQQSGEAEALARVYSSSISNGLSNLN NETSGTYANGSVIDLPKSEGYYNVVSGQPSP DQSGLDMT\GIKQIKQEPIYDLTSVPNLFTY\SS FNN\GQLAPGIT\MTEIDRIAQNIIKSHLETCQY TMEELHQLAWQTHTYEEIKAYQSKSREALW QQCAIQITHAIQYVVEFAKRITGFMELCQNDQ ILLLKSGCLEVVLVRMCRAFNPLNNTVLFEG KYGGMQMFKALGSDDLVNEAFDFAKNLCSL QLTEEEIALFSSAVLISPDRAWLIEPRKVQKLQ EKIYFALQHVIQKNHLDDETLAKLIAKIPTITA VCNLHGEKLQVFKQSHPEIVNTLFPPLYKELF NPDCATACK
792	2142	A	6440	92	781	SRGTFRCFCRDFFPCFSNMRLFLWNAVLTLFV TSLIGALIPEPEVKIEVLQKPFICHRKTKGGDL MLVHYEGYLEKDGSLFHSTHKHNNGQPIWFT LGILEALKGWGPGA*K/DMCVGEKRKLIIPPA LGYGKEGKGKIPPESTLIFNIDLLEIRNGPRSH ESFQEMDLNDDWKLSKDEVKAYLKKEFEKH GAVVNESHHDALVEDIFDKEDEDKDGFISAR EFTYKHDEL
793	2143	A			152	PRLKRLVVTEEDGGARPEALGKIAPRTPAELG ARADQELVTALMCDLRRPAAGGMMDLAYV CEWEK WSKSTHCPSVPLACAWSCRNLIAFTM DLRSDDQDLTRMIHILDTEHPWDLHSIPSEHH EAITC\LEWDQSGFPGFLFSRWPTGQIK\CWS MGVSTLA\NSWE\SSVGSL\VEGGPHLWALS\ WLH\NGVKLALHVEKSGASSFGEKFSR\VKFS P\SLTLF\GGNAMEGWIAVTVSGLVTVSLLQ\P SGQVL\TST\ESLCRLRARVALADIAFTGGGNI VVATADGSSA\SPVQFYKVCVSVVSEKCRIDT DILPSLFMRCTTDLNRKDKFPAITHLKFLARD MSEQVLLCASSQTSSIVECWSLRKEGLPVNNI FQQISPVVGDKQPTILKWRILSATNDLDRVSA V\ALPKLPISLTNTDLKVASDTQFYPGLGLAL AFHDGSVHIVHRLSLQTMAVFYSSAAPRVD EPAMKRPRTAGPAVHLKAMQLSWTSLALVG

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  IDSHGKLSVLRLSPSMGHPLEVGLALRHLLFL LEYCMVTGYDWWDILLHVQPSMVQSLVEKL HEEYTRQTAALQQVLSTRILAMKASLCKLSP CTVTRVCDYHTKLFLIAISSTLKSLLRPHFLNT PDKSPGDRLTEICTKITDVDIDKVMINLKTEEF VLDMNTLQALQQLLQWVGDFVLYLLASLPN QPCPTSEPCPTSEPSPTSEPSPTSEPSSP*SLC\G SLLRPGHSFLRDGTSLGMLRELMVVIRIWGLL KPSCLPVYTATSDTQDSMSLLFRLLTKLWICC RDEGPASEPDEALVDECCLLPSQLLIPSLDWL PASDGLVSRLQPKQPLRLQFGRAPTLPGSAAT LQLDGLARAPGQPKIDHLRRLHLGACPTEEC KACTRCGCVTMLKSPNRTTAVKQWEQRWIK NC/LVRWALVAGAPQLPLSPAAPQLLLSYPSA APEPGCCKSHRSPWTLLGAVNLSPPCRAVEG RGPDACVTSRASEEAPAFVQLGPQSTHHSPRT PRSLDHLHPEDRP
794	2144	A	6490	418	585	NGDKADLENESCRAQVLMPVVPALWEAEGG
795	2145	A	6499	395	1027	GSIEPRDLRLQ*AVITPL\TPAWVTQ KLLWLPPHSEQKRSPLYHPQGPSGTTPSAP\FS SHSPPPSLLQA\PSIAAFLRTHGHISASGPLRMP FPH/H*NAFLLVFPGQRSQLTS/PSHYLCREVFP DHHHHLCRLSLESSPLFHHRVLFCVPKQNVN STRAQIFCLFVHIVGCRCINTFPLHLFRLHLWL HFLQIPLCKKNKSVKLGKTVVGRGCQSAAGS DTRVRAAVGAPGLPVEPLV
796	2146	A	6503	68	936	HSALLTHSSFCVFTLCQDFFTYSSMSEEVTYA DLQFQNSSEMEKIPEIGKFGEKAPPAPSHVWR PAALFLTLLCLLLLIGLGVLASMFHVTLKIEM KKMNKLQNISEELQRNISLQLMSNMNISNKIR NLSTTLQTIATKLCRELYSKEQEHKCKPCPRR WIWHKDSCYFLSDDVQTWQESKMACAAQN ASLLKINNKNALEFIKSQSRSYDYWLGLSPEE DS/YSWYESG*YNQ\PSAWVIRNAPDLNNMY CGYINRLYVQYYHCTYKQRMICEKMANPVQ LGSTYFREA
797	2147	A	6507	1	881	PGSTHASARSQVPRSAGEAAPHSRRPPGLLPH APRAASAQLEERMRDPHPGMTLQEGDCRGS QTVSLTMGTADSDEMAPEAPQHTHIDVHIHQ ESALAKLLLTCCSALRPRATQARGSSRLLVAS WVMQIVLGILSAVLGGFFYIRDYTLLVTSGA AIWTGAVAVLAGAAAFIYEKRGGTYWALLR TLLALAAFSTAIAALKLWNEDFRYGYSYYNS ACRISSSSDWNTPAPTQSPEEVRRLHLCTSFM DMLKALFRTLQAMLLGVWILLLASLTPLWL /SL/RGECSQPKG*VPKKRDQKEMLEVSGI*PG STHASARSQVPRSAGEAAPHSRRPPGLLPHAP RAASAQLEERMRDPHPGMTLQEGDCRGSQT VSLTMGTADSDEMAPEAPQHTHIDVHIHQES ALAKLLTCCSALRPRATQARGSSRLLVASW VMQIVLGILSAVLGGFFYIRDYTLLVTSGAAI WTGAVAVLAGAAAFIYEKRGGTYWALLRTL LALAAFSTAIAALKLWNEDFRYGYSYYNSAC RISSSSDWNTPAPTQSPEEVRRLHLCTSFMDM LKALFRTLQAMLLGVWILLLASLTPLWLYC WRMFPTKGVSP
798	2148	A	6528	912	2287	VPNYLPSVSSAIGGEVPQRYVWRFCIGLHSAP RFLVAFAYWNHYLSCTSPCSCYRPLCRLNFG LNVVENLALLVLTYVSSSEDF/TWVPG*GRSG

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  EVFPEGTGLPLPHSDLPTSWCGHSLQCGSQSS FPPAIHENAFIVFIASSLGHMLLTCILWRLTKK HTVSQE\DGLSLAGAPRQPRRKSRTSVLRIRV MVRWELSSNGNPGRGVLGLGLGLGNKLRVV GQNLGL*HCVWVVWETGE*KRWRLQMGIE*GVASRRQ*VRNSVRGLVCHNSSAPPMYMGFF SPTVFGGGVGG*LHVTFILHPPEVEAAGIPLLL GPSLPQRQGREHIVVILAAPACAPFHDR*WEP REIRPSP*ELGLRGEPTLSYPASCRVIRQPIP*D RKSYSWKQRLFIINFISFFSALAVYFRHNMYC EAGVYTIFAILEYTVVLTNMAFHMTAWWDF
799	2149	A	6529	1	874	GNKELLITSQPEEKRF FFFFQRINPIEHSGSVSLLALACDLGWCEDWS CCLVQGGGDLVDVVQTNHGEDEAGGDTDSV DEARCKESQQEAQENLREDLCLESFAKDKIL QIIEGSEREHEETRTKQAALDGEPLGGGQLTA VHLHPSKEQQGQEGGERQRGARTHHWRGW EKGRRVRLRPPSGKLRADQPVRKLGGPTPS/T ELPGLQPHAPTPHTA/PATPTYSPAPDTPNPPV RWKCPLPVEPRTRQLCRERTRKACPPKPRPPL GLPGDPTGPVTHHAPPVSPTGASGQERRAEP GAVSYAHASATK
800	2150	A	6544	2		SAQRWAAVAGRWGCRLLALLLLVPGPGGAS EITFELPDNAKQCFYEDIAQGTKCTLEFQVITG GHYDVDCRLEDPDGKVLYKEMKKQYDSFTF TASKNGTYKFCFSNE\FSTFTHKTVYFDFQVG E\THLCFLVR/DRVSALTQMESACVSIHEALKS VIDYQTHFRLREAQGRSRAEDLNTRVAYWSV GEALILLVVSIGQVFLLKSFFSDKRTTTTRVGS
801	2151	A .	6556	1	1319	TPCMECIKGEGLREPQNLSGSQREPQTEGSM DGWRRMPRWGLLLLLWGSCTFGLPTDTTTF KRIFLKRMPSIRESLKERGVDMARLGPEWSQP MKRLTLGNTTSSVILTNYMDTQYYGEIGIGTP PQTFKVVFDTGSSNVWVPSSKCSRLYTACVY HKLFDASDSSSYKHNGTELTLRYSTGTVSGFL SQDIITVGGITVTQMFGEVTEMPALPFMLAEF DGVVGMGFIEQAIGRVTPIFDNIISQGVLKED VFSFYYNRDSENSQSLGGQIVLGGSDPQHYE GNFHYINLIKTGVWQIQMKGVSVGSSTLLCE DGCLALVDTGASYISGSTSSIEKLMEALGAKE KRLFDYVVKCNEGPTLPPTFLFLLGGKDTPLT SADYLFQESYSSKKLSTLAIHAMYIPPTGPTL \ALGATF\IRKFYTEFDRGNNPHGFALAR
802	2152	A	6567	13	6147	MCLGRMGASSPRSPEPVGPPAPGLPFCCGGSL LAVVVLLALPVAWGQCNAPEW\LPFARPTNL TDEFEFPIGTYLNYECRPGYSGRPFSIICLKNS VWTGAKDRCRRKSCRNPPDPVNGMVHVIKG IQFGSQIKYSCTKGYRLIGSSSATCIISGDTVIW DNETPICDRIPCGLPPTITNGDFISTNRENFHY GSVVTYRCNPGSGGRKVFELVGEPSIYCTSND DQVGIWSGPAPQCIIPNKCTPPNVENGILVSD NRSLFSLNEVVEFRCQPGFVMKGPRVKCQA LNKWEPELPSCSRVCQPPPDVLHAERTQRDK DNFSPGQEVFYSCEPGYDLRGAASMRCTPQG DWSPAAPTCEVKSCDDFMGQLLNGRVLFPV NLQLGAKVDFVCDEGFQLKGSSASYCVLAG MESLWNSSVPVCEQIFCPSPPVIPNGRHTGKP LEVFPFGKAVNYTCDPHPDRGTSFDLIGESTIR CTSDPQGNGVWSSPAPRCGILGHCQAPDHFL FAKLKTQTNASDFPIGTSLKYECRPEYYGRPF

	icted end Amino acid sequence (A=Alanine C=Cysteine,
	eotide D=Aspartic Acid, E=Glutamic Acid,
nucl- peptide in nucleotide locat	
eotide seq- USSN location corre	sponding   I=Isoleucine, K=Lysine, L=Leucine,
seq- uence 09/496 correspondi to las	st amino M=Methionine, N=Asparagine, P=Proline,
	residue Q=Glutamine, R=Arginine, S=Serine,
	reptide T=Threonine, V=Valine, W=Tryptophan,
residue of seque	
peptide	/=possible nucleotide deletion, \=possible
sequence	nucleotide insertion
	SITCLDNLVWSSPKDVCKRKSCKTPPDPVNG
	MVHVITDIQVGSRINYSCTTGHRLIGHSSAECI -
	LSGNAAHWSTKPPICQRIPCGLPPTIANGDFIS
	TNRENFHYGSVVTYRCNPGSGGRKVFELVGE
	PSIYCTSNDDQVGIWSGPAPQCIIPNKCTPPNV
	ENGILVSDNRSLFSLNEVVEFRCQPGFVMKGP
	RRVKCQALNKWEPELPSCSRVCQPPPDVLHA
	· ERTQRDKDNFSPGQEVFYSCEPGYDLRGAAS
	MRCTPQGDWSPAAPTCEVKSCDDFMGQLLN
	, ,
	GRVLFPVNLQLGAKVDFVCDEGFQLKGSSAS
	YCVLAGMESLWNSSVPVCEQIFCPSPPVIPNG
	RHTGKPLEVFPFGKAVNYTCDPHPDRGTSFD
	LIGESTIRCTSDPQGNGVWSSPAPRCGILGHC
	QAPDHFLFAKLKTQTNASDFPIGTSLKYECRP
	EYYGRPFSITCLDNLVWSSPKDVCKRKSCKTP
	PDPVNGMVHVITDIQVGSRINYSCTTGHRLIG
	HSSAECILSGNTAHWSTKPPICQRIPCGLPPTI
	ANGDFISTNRENFHYGSVVTYRCNLGSRGRK
	VFELVGEPSIYCTSNDDQVGIWSGPAPQCIIPN
	KCTPPNVENGILVSDNRSLFSLNEVVEFRCOP
	GFVMKGPRRVKCQALNKWEPELPSCSRVCQ
	PPPEILHGEHTPSHQDNFSPGQEVFYSCEPGY
	DLRGAASLHCTPQGDWSPEAPRCAVKSCDDF
	LGQLPHGRVLFPLNLQLGAKVSFVCDEGFRL
	KGSSVSHCVLVGMRSLWNNSVPVCEHIFCPN
	PPAILNGRHTGTPSGDIPYGKEISYTCDPHPDR
	GMTFNLIGESTIRCTSDPHGNGVWSSPAPRCE
	LSVRAGHCKTPEQFPFASPTIPINDFEFPVGTS
	LNYECRPGYFGKMFSISCLENLVWSSVEDNC
	RRKSCGPPPEPFNGMVHINTDTQFGSTVNYSC
	NEGFRLIGSPSTTCLVSGNNVTWDKKAPICEII
	SCEPPPTISNGDFYSNNRTSFHNGTVVTYQCH
	TGPDGEQLFELVGERSIYCTSKDDQVGVWSS
	PPPRCISTNKCTAPEVENAIRVPGNRSFFSLTEI
	IRFRCQPGFVMVGSHTVQCQTNGRWGPKLPH
	CSRVCQPPPEILHGEHTLSHQDNFSPGQEVFY
	SCEPSYDLRGAASLHCTPQGDWSPEAPRCTV
	KSCDDFLGQLPHGRVLLPLNLQLGAKVSFVC
	DEGFRLKGRSASHCVLAGMKALWNSSVPVC
	EQIFCPNPPAILNGRHTGTPLGDIPYGKEVSYT
	CDPHPDRGMTFNLIGESTIRRTSEPHGNGVWS
	SPAPRCELPVGAACPHPPKIQNGHYIGGHVSL
	,
	YLPGMTISYTCDPGYLLVGKGFIFCTDQGIWS
	QLDHYCKEVNCSFPLFMNGISKELEMKKVYH
	YGDYVTLKCEDGYTLEGSPWSQCQADDRWD
	PPLAKCTSRTHDALIVGTLSGTIFFILLIIFLSWI
	ILKHRKGNNAHENPKEVAIHLHSQGGSSVHP
	RTLQTNEENSRVLP
803 2153 A 6574 2 3233	HGRSARLAAVPAEAMPGPRRPAGSRLRLLLL
	LLLPPLLLLRG\SHAGNLTVAVVLPLANTSY
	PWSWA\RVGPAVELALAQVKARPDLLPGWT
	VRTVLGSSENALGVCSDTAAPLAAVDLKWF
	HNPAVFLGPGCVYAAAPVGRFTAHWRVPLL
	TAGAPALGFGVKDEYALTTRAGPSYAKLGDF
	VAALHRRLGWERQALMLYAYRPGDEEHCFF
	LVEGLFMRVRDRLNITVDHLEFAEDDLSHYT
	RLLRTMPRKGRVIYICSSPDAFRTLMLLALEA
	GLCGEDYVFFHLDIFGQSLQGGQGPAPRRPW
	ODCODD I VIXIDDII OQUDOGOOTII KKI W
	ERGDGQDVSARQAFQAAKIITYKDPDNPEYL EFLKQLKHLAYEQFNFTMEDGLVNTIPASFH

COOLD	CEO TE	N.4-+	LODA	I D., 41-2-1	1 D. 31.2	A !
SEQ ID NO: of	SEQ ID NO: of	Met hod	SEQ ID NO:	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide	2200	in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence	ł	09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence			914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
				amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
ł	l	ł	1	residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
	ļ			peptide sequence		/=possible nucleotide deletion, \=possible nucleotide insertion
<u> </u>	<del></del>	<del> </del>		sequence		DGLLLYIQAVTETLAHGGTVTDGENITQRMW
	[	[	[			NRSFQGVTGYLKIDSSGDRETDFSLWDMDPE
						NGAFRVVLNYNGTSQELVAVSGRKLNWPLG
		]		J	J	YPPPDIPKCGFDNEDPACNODHLSTLEVLALV
						GSLSLLGILIVSFFIYRKMQLEKELASELWRVR
					·	WEDVEPSSLERHLRSAGSRLTLSGRGSNYGSL
l	1					LTTEGQFQVFAKTAYYKGNLVAVKRVNRKR
						IELTRKVLFELKHMRDVQNEHLTRFVGACTD
	[					PPNICILTEYCPRGSLQDILENESITLDWMFRY
						SLTNDIVKGMLFLHNGAICSHGNLKSSNCVV DGRFVLKITDYGLESFRDLDPEQGHTVYAKK
		ļ	}			LWTAPELLRMASPPVRGSQAGDVYSFGIILQE
						IALRSGVFHVEGLDLSPKEIIERVTRGEOPPFR
		l				PSLALQSHLEELGLLMQRCWAEDPQERPPFQ
	1		1			QIRLTLRKFNRENSSNILDNLLSRMEQYANNL
						EELVEERTQAYLEEKRKAEALLYQILPHSVAE
						QLKRGETVQAEAFDSVTIYFSDIVGFTALSAE
Ì	ĺ	1				STPMQVVTLLNDLYTCFDAVIDNFDVYKVET
		1				IGDAYMVVSGLPVRNGRLHACEVARMALAL
	]					LDAVRSFRIRHRPQEQLRLRIGIHTGPVCAGV VGLKMPRYCLFGDTVNTASRMESNGEAL\KI
						HLSS\ETKAVL\EEFGGFELELRGDVEMKGKG
						KVRTYWLLGERGSSTRG
804	2154	A	6585	2	3837	DAPGRPPVRLPTMELEDGVVYQEEPGGSGAV
						MSERVSGLAGSIYREFERLIVRYDEEVVKELIP
ĺ						LVVAVLENLDSVFAQDQEHQVELELLRDDNE
	Ì					QLITQYEREKALRKHAEEKFIEFEDSQEQEKK
						DLQTRVESLESQTRQLELKAKNYADQISILEE
						REAELKKEYNALHQRHTEMIHNYMEHLERT KLHQLSGSDQLESTAHSRIRKERPISLGIFPLP
					ı	AGDGLLTPDAQKGGETPGSEQWKFQELSQPR
						SHTSLKDELSDVSQGGSKATTPASTANSDVA
			ļ ,	,		TIPTDTPLKEENEGFVKVTDAPNKSEISKHIEV
				!		QVAQETRNVSTGSAENEEKSEVQAIIESTPEL
			1			DMDKDLSGYKGSSTPTKGIENKAFDRNTESL
						FEELSSAGSGLIGDVDEGADLLGMGREVENLI
			]			LENTQLLETKNALNIVKNDLIAKVDELTCEK
						DVLQGELEAVKQAKLKLEEKNRELEEELRKA RAEAEDARQKAKDDDDSDIPTAQRKRFTRVE
						MARVLMERNQYKERLMELQEAVRWTEMIR
						ASRENPAMQEKKRSSIWQFFSRLFSSSSNTTK
						KPEPPVNLKYNAPTSHVTPSVKKRSSTLSQLP
						GDKSKAFDFLSEETEASLASRREQKREQYRQ
						VKAHVQKEDGRVQAFGWSLPQKYKQVTNG
						QGENKMKNLPVPVYLRPLDEKDTSMKLWCA
						VGVNLSGGKTRDGGSVVGASVFYKDVAGLD
						TEGSKQRSASQSSLDKLDQELKEQQKELKNQ EELSSLVWICTSTHSATKVLIIDAVQPGNILDS
						FTVCNSHVLCIASVPGARETDYPAGEDLSESG
						OVDKASLCGSMTSNSSAETDSLLGGITVVGC
				ļ		SAEGVTGAATSPSTNGASPVMDKPPEMEAEN
						SEVDENVPTAEE\ATEATEGNAGSAEDTV\DIS
					l	QTGVYTEHVFTDPLG\VQIPEDLSPVYQSSND
				l		SDAYKDQISVLPNEQDLVREEAQKMSSLLPT
				l		MWLGAQNGCLYVHSSVAQWRKCLHSIKLKD
			(	ľ	i	SILSIVHVKGIVLVALADGTLAIFHRGVDGQW
			]	}	j	DLSNYHLLDLGRPHHSIRCMTVVHDKVWCG
						YRNKIYVVQPKAMKIEKSFDAHPRKESQVRQ LAWVGDGVWVSIRLDSTLRLYHAHTYQHLO
						DVDIEPYVSKMLGTGKLGFSFVRITALMVSC
			L			

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  NRLWVGTGNGVIISIPLTETVILHQGRLLGLR  ANKTSGVPGNRPGSVIRVYGDENSDKVTPGT  FIPYCSMAHAQLCFHGHRDAVKFFVAVPGQV  ISPQSSSSGTDLTGDKGRGHLHRSLVVRRP
805	2155	A	6605	469	2602	FGRLLWGTAFKSWKMKAPPHLILLYATFTQ SLKVVTKRGSADGCTDWSIDIKKYQVLVGEP VRIKCALFYGYIRTNYSLAQSAGLSLMWYKS SGPGDFEEPIAFDGSRMSKEEDSIWFRPTLLQ DSGLYACVIRNSTYCMKVSISLTVGENDTGL CYNSKMKYFEKAELSKSKEISCRDIEDFLLPT REPEILWYKECRTKTWRPSIVFKRDITLIREV REDDIGNYTCELKYGGFVVRRTTELTVTAPL TDKPPKLLYPMESKLTIQETQLGDSANLTCRA FFGYSGDVSPLIYWMKGEKFIEDLDENRVWE SDIKILKEHLGEQEVSISLIVDSVEEGDLGNYS CYVENGNGRRHASVLLHKRELMYTVELAGG LGAILLLLVCLVTIYKCYKIEIMLFYRNHFGA EELDGDNKDYDAYLSYTKVDPDQWNQETGE EERFALEILPDMLEKHYGYKLFIPDRDLIPTGT YIEDVARCVDQSKRLIIVMTPNYVVRRGWSIF ELETRLRNMLVTGEIKVILIECSELRGIMNYQE VEALKHTIKLLTVIK WHGPKCNKLNSKFWKR LQYEMPFKRIEPITHEQALDVSEQGFFGELQT VSAISMAAATSTALATAHPDLRSTFHNTYHS QMRQKHYYRSYEYDVPPTGTLPLTSIGNQHT YCNIPMTLINGQRPQTKSSREQNPDEAHTNSA ILPLLPRETSISSVIW
806	2156	A	6614	3	1584	NSARGGVGVRGARAMATVQEKAAALNLSAL HSPAHRPPGFSVAQKPFGATYVWSSIINTLQT QVEVKKRRHRLKRHNDCFVGSEAVDVIFSHL IQNKYFGDVDIPRAKVVRVCQALMDYKVFE AVPTKVFGKDKKPTFEDSSCSLYRFTTIPNQD SQLGKENKLYSPARYADALFKSSDIRSASLED LWENLSLKPANSPHVNISTTLSPQVINEVWQE ETIGRLLQLVDLPLLDSLLKQQEAVPKIPQPK RQSTMVNSSNYLDRGILKAYSDSQEDEWLSA AIDCLEYLPDQMVVEISRSFPEQPDRTDLVKE LLFDAIGRYYSSREPLLNHLSDVHNGIAELLV NGKTEIALEATQLLLKLLDFQNREEFRRLLYF MAVAANPSEFKLQKESDNRMVVKRIFSKAIV DNKNLSKGKTDLLVLFL\MDHQKDVFKIPGT L\HKIVS\VK\LMAIQNGRDPNRDAGYIYCQRI DQRDYSNITEKTTIDELLYLLKTLDEDSKLSA KEKKK\LLGQFYKCHPDIFIEHFGD
807	2157	A	6615	4198	2094	FGIVGTFALETDELDSDRDPAIFSLCDFGAMR PQILLLALLTLGLAAQHQDKVPCKM/VKML CPDRVDKKVSCQVLGLLQVPSVLPPDTETLD LSGNQLRSILASPLGFYTALRHLDLSTNEISFL QPGAFQALTHLEHLSLAHNRLAMATALSAG GLGPLPRVTSLDLSGNSLYSGLLERLLGEAPS LHTLSLAENSLTRLTRHTFRDMPALEQLDLHS NVLMDIEDGAFEGLPRLTHLNLSRNSLTCISD FSLQQLRVLDLSCNSIEAFQTAS\QPQAEFQLT WLDLRENKLLHFPDLAALPRLIYLNLSNNLIR LPTGPPQDSKGIHAPSEGWSALPLS\APSGNAS GRPLSQLLNLDLSYNEIELIPDSFLEHLTSLCFL NLSRNCLRTFEARRLGSLPCLMLLDLSHNALE TLELGARALG\SLRTLLLQGNALRDLPPYTFA NLASLQRLNLQGNRVSPCGGPDEPGP\SGCV\ AFSGITSLRSLSLVDNEIELLRAGAFLHTPLTE

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isolcucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  LDLSSNPGLEVATGALGGLEASLEVLALQGN GLMVLQVDLPCFICLKRLNLAENRLSHLPGN TO AVEL EVIL DELEGALGGLEASLEV DELEGALGGLEASURE DELEG
					5	TQAVSLEVLDLRNNSFSLLPGSAMGGLETSLR RLYLQGNPLSCCGNGWLAAQLHQGRVDVDA TQDLICRFSSQEEVSLSHVRPEDCEKGGLKNI NLIIILTFILVSAILLTTLAACCCVRRQKFNQQ YKA
808	2158	A	6619	153	1852	FKALSQYIYTNTHLEREAAFEVAILLRRMEEG ARHRNNTEKKHPGGGESDASPEAGSGGGGV ALKKEIGLVSACGIIVGNIIGSGIFVSPKGVLEN AGSVGLALIVWIVTGFITVVGALCYAELGVNI PKSGGDYFYVKDIFGGLAGFLRLWIAVLVIYP TNQAVIALTFSNYVLQPLFPTCFPPESGLRLLA AICLLLLTWVNCSSVRWATRVQDIFTAGKLL ALALIIIMGIVQICKGEYFWLEPKNAFENFQEP DIGLVALAFLQGSFAYGGWNFLNYVTEELV DPYKNL\PRAIFISIP\LVTFVYVFANV/ALYVT AMSPQEL\LAS\NAVAVTFGEKLLGVMAWIM PISVALSTFGGVNGSLFTSSRLFFAGAREGHLP SVLAMIHVKRCTPIPALLFTCISTLLMLVTSD MYTLINYVGFINYLFYGVTVAGQIVLRWKKP DIPRPIKINLLFPIIYLLFWAFLLVFSLWSEPVV CGIGLAIMLTGVPVYFLGVYWQHKPKCFSDFI ELLTLVSQKMCVVVYPEVERGSGTEEANED MEEQQQPMYQPTPTKDKDVAGQVPQP
809	2159	A	6621	1041	223	QDSRKMLPSTSVNSLVQGNGVLNSRDAARH TAGAKRYKYLRRLFRFRQMDFEFAAWQMLY LFTSPQRVYRNFHYRKQTKDQWARDDPAFL VLLSIWLCVSTIGFGFVLDMGFFETIKLLLWV VLIDCVGVGLLIATLMWFISNKYLVKRQSRD YDVEWGYAFDVHLNAFYPLLVILHFIQLFFIN HVILTDTFIGYLVGNTLWLVAVGYYIYVTFL GYSVGLLFFS\ALPFLKNTVILLYPFAPLILLYG LSLALGWNFTHTLCSFYKYRVK
810	2160	A	6623	160	822	SPASGHCRLNGAAVAMFGCLVAGRLVQTAA QQVAEDKFVFDLPDYESINHVVVFMLGTIPFP EGMGGSVYFSYPDSNGMPVWQLLGFVTNGK PSAIFKISGLKSGEGSQHPFGAMNIVRTPSVAQ IGISVELLDSMAQQTPVGNAAVSSVDSFTQFT QKMLDNFYNFASSFAVSQ/VPDDTQ/RPSEMF IPANVVLKWYENFQRRTSTEPSLLENIIWIKIN F
811		A	6627	18	3367	LEGSLNTERAKYYLTITMPHFTVTKVEDPEEG AAASISQEPSLADIKARIQDSDEPDLSQNSITG EHSQLLDDGHKKARNAYLNNSNYEEGDEYF DKNLALFEEEMDTRPKVSSLLNRMANYTNLT QGAKEHEEAENITEGKKKPTKTPQMGTFMG VYLPCLQNIFGVILFLRLTWVVGTAGVLQAF AIVLICCCCTMLTAISMSAIATNGVVPAGGSY FMISRALGPEFGGAVGLCFYLGTTFAAAMYIL GAIEIFLVYIVPRAAIFHSDDALKESAAMLNN MRVYGTAFLVLMVLVVFIGVRYVNKFASLFL ACVIVSILAIYAGAIKSSFAPPHFPVCMLGNRT LSSRHIDVCSKTKEINNMTVPSKLWGFFCNSS QFFNATCDEYFVHNNVTSIQGIPGLASGIITEN LWSNYLPKGEIIEKPSAKSSDVLGSLNHEYVL VDITTSFTLLVGIFFPSVTGIMAGSNRSGDLKD AQKSIPIGTILAILTTSFVYLSNVVLFGACIEGV VLRDKFGDAVKGNLVVGTLSWPSPWVIVIGS FFSTCGAGLQSLTGAPRLLQAIAKDNIIPFLRV

SEQ ID NO: of nucl- cotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
						FGHSKANGEPTWALLLTAAIAELGILIASLDL VAPILSMFFLMCYLFVNLACALQTLLRTPNW RPRFRYYHWALSFMGMSICLALMFISSWYYA IVAMVIAGMIYKYIEYQGAEKEWGDGIRGLS LSAARFALLRLEGPPHTKNWRPQLLVLLKL DEDLHVKHPRLLTFASQLKAGKGLTIVGSVIV GNFLENYGEALAAEQTIKHLMEAEKVKGFCQ LVVAAKLREGISHLIQSCGLGGMKHNTVVM GWPNGWRQSEDARAWKTFIGTVRVTTAAHL ALLVAKNISFFPSNVEQFSEGNIDVWWIVHDG GMLMLLPFLLK\QHKVWRKCSIRFFYTVAQLE DNSIQMKKDLATFLYHLRIEAEVEVVEMHDS DISAYTYERTLMMEQRSQMLRHMRLSKTER DREAQLVKDRNSMLRLTSIGSDEDEETETYQ EKVHMTWTKDKYMASRGQKAKSMEGFQDL LNMRPDQSNVRRMHTAVKLNEVIVNKSHEA KLVLLNMPGPPRNPEGDENYMEFLEVLTEGL ERVLLVRGGGSEVITIYS
812	2162	A	6628	66	640	AVCTMSEMAELSELYEESSDLQMDVMPGEG DLPQMEVGSGSRELSLRPSRSGAQQLEEEGP MEEEEAQPMAAPEGKRSLANGPNAGEQPGQ VAGADFESEDEGEEFDDWEDDYDYPEEEQLS GAGYRVSAALEEADKMFLRTREPALDGGFQ MHYEKTPFDQLAFIEELF\SLMVVNRLTEELG CDEIIDRE
813	2163	A	6630	708	1355	AKMGAYKYIQELWRKKQSDVMRFLLRVRC WQYRQLSALHRAPRPTRPDKARRLGYKAKQ GY/VYIYIGFVFAVIYRIRVRRGGRKRPVPKG ATYGKPVHHGVNQLKFARSLQSVAEERAGR HCGALRVLNSYWVGEDSTYKFFEVILIDPFHK AIRRNPDTQWITKPVHKHREMRGLTSAGRKS RGLGKGHKFHHTIGGSRRAAWRRRNTLQLH RYR
814	2164	A	6635	201	1705	KGTEMNKSRWQSRRRHGRRSHQQNPWFRLR DSEDRSDSRAAQPAHDSGHGDDESPSTSSGT AGTSSVPELPGFYFDPEKKRYFRLLPGHNNCN PLTKESIRQKEMESKRLRLLQEEDRRKKIARM GFNASSMLRKSQLGFLNVTNYCHLAHELRLS CMERKKVQIRSMDPSALASDRFNLILADTNS DRLFTVNDVTVGGSKYGIINLQSLKTPTLKVF MHENLYFTNRKV\NSVCWASLNHLDSHILLC LMGLAETPGCATLLPASLFVNSHPAGIDRPG\ MLCSFRIPGAWSCAWSLNIQANNCFSTGLSR RVLLTNVVTGHRQSFGTNSDVLAQQFALMA PLLFNGCRSGEIFAIDLRCGNQGKGWKATRLF HDSAVTSVRILQDEQYLMASDMAGKIKLWD LRTTKCVRQYEGHVNEYAYLPLHVHEEEGIL VAVGQDCYTRIWSLHDARLLRTIPSPYPASKA DIPSVAFSSRLGGSRGAPGLLMAVGQDLYCY SYS
815	2165	A	6643	659	3282	NKNILEVPSARTTRIMGDHLDLLLGVVLMAG PVFGIPSCSFDGRIAFYRFCNLTQVPQVLNTTE RLLLSFNYIRTVTASSFPFLEQLQLLELGSQYT PLTIDKEAFRNLPNLRILDLGSSKIYFLHPDAF QGLFHLFELRLYFCGLSDAVLKDGYFRNLKA LTRLDLSKNQIRSLYLHPSFGKLNSLKSIDFSS NQIFLVCEHELEPLQGKTLSFFSLAANSLYSR VSVDWGKCMNPFRNMVLEILDVSGNGWTV DITGNFSNAISKSQAFSLILAHHIMGAGFGFHN IKDPDQNTFAGLARSSVRHLDLSHGFVFSLNS

TPSENPSLEQLFLGERMLQLAWETELCWD BEISHLQVI_NLNINVINSIPPOVIPSHLTAI GLSI.NSNRLTVLSHDLPANLEILDISRNQI APNPDVPSU.SVJ.DITHNCFCCELS.FTIN HTINVTIAOPPADIYCVYPDSLSGVSL.FSL. GCDEEPVI_SCL_WELTHNLTILTIJAITI TKPEGFCTCYKT.AQRLVFKDHPQGTEPDM KYDAYLCFSSKDFTWVQNALLKHDTQY. QNRPNLCFEERDFVPGENRPANNQDAIWM KIVCLVSRIFIRDGWCLEAFSYAQRCLS. NSALIMVYVGSLSQYQLMKHQSIRGFVQK YLRWPEDLQDVGWFLHKL.SQQILKKEKE KONNIPLIQTVATIS  816 2166 A 6646 I 3811 RDRAGVRPAGKQHAAAAAFYDVGGRDY GGARLASLFGLDQAAAGHGNEFFOYTAPK KKQQTAATAGNQATPKTMFLEMVRTM STYDDQRQMVWSIMFBSEKAAVEPNKQV KKQQTAATAGNQATPKTMFLEMVRTM STYYDDQRQMVWSIMFBSEKAAVEPNKQV KCNSTSSLDAVLSQDLIVMFLEMVRTM STYYDDQRQMVWSIMFBSEKAAVEPNKQV KCNSTSSLDAVLSQDLIVMFLEMVRTM STYYDDQRQMVWSIMFBSEKAAVEPNKQV KCNSTSSLDAVLSQDLIVMFLEMVRTM STYYDDQRQMVWSIMFBSEKAAVEPNKQV KCNSTSSLDAVLSQDLIVMFLEMVRTM STYYDDQRQMVWSIMFBSEKAAVEPNKQV KCNSTSSLDAVLSQDLIVMFLEMVRTM STYYDDQRQMVWSIMFBSEKAAVEPNKQV KCNSTSSLDAVLSQDLIVMFLEMVRTM STYYDDQRQMVWSIMFBSEKAAVEPNKQV KCNSTSSLDAVLSQDLIVMFLEMVRTM STYYDDQRQMVWSIMFBSEKAAVEPNKQV KCNSTSSLDAVLSQDLIVMFLEMVRTM STYPDDQRQMVTSVBLQAVATABSTPILL GSKLVIKGWEDGML.GMCGGRLLIVM CAVAKAKLISMAKMGQPALPILPPQLDSND EDVNTLQGGQGPVTTFSVQPALVEND STYPLDDQRQGPVTFSVQSLQAAHAAL MTSQAPQPSVTGLQAPSAALMQVSSLDSH VSGNAQSFQPYAGMQAYAYQASAVTSQ PVRRLYPAPLSQPPHFQGSGDMASFLMTE QMUTESLKAMVSKVADKMBKILLERNQRYVE NLMMEKRNNSLQTATENTQARVLHACQE KVTBELAAATAQVSHLQLMVTAHQKE TVRELAAATAQVSHLQLMVTAHQKE TVRELAAATAQVSHLQLMVTAHQKE TVRELAAATAQVSHLQLMVTAHQKE TVRELAAATAQVSHLQLKVTSLEELI RVEKESLEKNLSERKKKSQDRSQAGEED RKSVQEELDKLRQLKKTRVSTDQAAABC LVQAGLQTQWFAKCCHLLASAKDEHLQQ EVCAQRDAYQQKLVQLQEKSESVECACLAL QTTALTKQNEQHILKELIKKNSQMSGVBAA DPSEKVKKIMNQVPGSLEKVTSLEELI RVEKESLEKNLSERKKKSAQERSQAGEED RKSVQEELDKLRQLKKTRVSTDQAAABC LVQAGLQTQWPAKCCHLLASAKDEHLQQ EVCAQRDAYQQKLVQLQEKESSTELESSYNGE EKAEERPRRPSQEQSAASSGQPQABANE ESSMVPSBQVVEEAVPLPQALTTSQDGHR KGSSEABALSEIKDGSLPFELLSESPNENG ESSMVPSBQVVEEAVPLPQALTTSQDGHR KGSSEABALSEIKDGSLPFELLSESPNENG ESSMVPSBQVVEEAVPLPQALTTSQDGHR ESSTEAGSTVAGAAALRRSHHSQRSSLSGD DELFKGATLKALRPKAQQFEQEEDEDSVSMK				11.			
nucleotide sequence control (USSN) control (1944) and the sequence							
cotide sequence were purposed by the sequence of the sequence	1		hod	ID NO:			
uence uence 914 got correspondi pol ast amino acid residue of peptide sequence peptide sequence peptide sequence peptide sequence peptide sequence peptide sequence peptide sequence peptide sequence peptide sequence peptide sequence peptide sequence peptide sequence peptide sequence peptide sequence peptide sequence peptide sequence peptide sequence peptide		peptide			1		
genice   g	eotide	seq-		USSN	location	corresponding	
amino acid residue of peptide sequence    T-Threonine, V=Valine, W=Tryptophan, y=cytosine, X=Unknown, *=Stop codon, y=posible nucleotide deletion, V=possible nucleotide insertion   RVFETLEDLEVENLAYNERIKADEAFYG   NQVVLN.SYNLLGELYSSNPYGEPKVAYII   QKNHIAIIQDQTFKFLEKLQTLDLENDNATT   FPSIPPIELSORIK, VTLFKINLTANLHLISE   LENLDILYFLRVPHQUILINONRESSCSG   TPSENFSLEQLE-GEPSIMIQLA WEBTELCWD   EGLSHLQVLVLN-SNNLTUSLEDISNOQ   APNPDVFVSLSVLDITHNIKFICECELSTEIN   NITHNYTIAGP-ADJY CVYPPSLS-GVS-ST-SL-   GCDEEFVLKSLKFS IFTVCTVILT-I-I-I-I-IIII   TFRGFCFICVKTAGRU-KVBAPPGTEPDN   KYDAYLCPSSKPFTW-QNALLKHLDTOY,   QREPHIC-PEERPYPGENRAN-AND-AND-AND-AND-AND-AND-AND-AND-AND-A	seq-	uence	1	09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
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residue of peptide sequence peptide sequ		}	l	1	amino acid	of peptide	
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RVPETI.KDLKVI.NIA.YNKINKIADE.AFYGO NICQUALIS.YNLIGELYSSNIYGJERVAYTI QKNHIAIGDQTFKFLERLQTI.DLR.DNALT FPISPIDIFLSGKI.VTLYENLAI.HLI.SEE ELENLDI.YFLLR.VPH.OJLII.NQNRFSSC.GG TPSEPSIE.GG.LFGENM.Q.G. WETELCWD EGI.SH.QVI.VI.NHNYI.NS.PPGVPSH.T.AI GLSI.NSNIL.TVLSSNID.PAB.ELI.DISI.NQ) APPPDVFVSLSV.DITHNKFICE.CEL.STIFT NHTINVTI.AGPPDJYCVY.PDS.SGVSI.FSL. GCDEEVI.KSI.KFSLFVCTYTI.T.I.FLATIII IT.KFRGFCFICE/KTIAQKLV.FWLOPGTEP.DN KYDAYL.CFSSKDFTW.VQNALI.KH.DTQY.V QNRRNI.CFEEBDFYPGENRF.NGDATWN KIVCLVSBHFLR.DGWCLEAFSYA.QGRGF.VQX YLRWFEDLQDVGWFLHKL.SQQLLKKEKE KDN.IPL.QTVATIS  SIALIMVVGSI.SQYQLMKKINGSIRGFVQX YLRWFEDLQDVGWFLHKL.SQQLLKKEKEE KDN.IPL.QTVATIS  SIALIMVVGSI.SQYQLMKKOSIRGFVQX YLRWFEDLQDVGWFLHKL.SQQLLKKEKEE KDN.IPL.QTVATIS  GGARLASLFGI.DQAAAGHGNFFFQYTSH GGARLASLFGI.DQAAAGHGNFFFQYTSH KKGGGTAATIONQATPKTAPATMSTPTIL.V AVHAYRYTNGQYVKQGKFGAAAULSHVITT EVRILLYISQQQVTVARIIIVNFELMVRIN STFYDDQRQNWSIMFESEKAAVEFNKQVV KCNSTSSL.DAVLSQDLLVAGGEFGAAAULSHVITT EVRILLYISQQVDVTVARIIVNFELMVRIN STFYDDQRQNWSIMFESEKAAVEFNKQVV KCNSTSSL.DAVLSQDLLVAGGERGAAVLGSHV KNSGSGGHISVSSRDSAAPSPIPGADNI.SAD VSFPTSIPFKSGEPALRTKSNSLSEQLAINT AVKAKLISRMAKMGQPM.DIVOSLDSH VSGNAGSFGYVGGMGRAYAYPQASAVTSQ BDVNTL.QGGQPVVTTSVQPSL.CPAHPAL MTSQAPGPSVTQL.QATSAALMSQRIMTE QHNTELRAAVSVADAKSMVARDHLMTKVEELQ SAGNSALIPSMSVTMETSMINGRIIQE LKQELLEKSNREEQNDXISELIERNGYSLMED PVPLYPAPLSQDFHGGGSGRAVINGRILIPP LKQELLEKSNREEQNDXISELIERNGYLAEQE KVTEELAAATAQVSHLQLKVTAHQKKKLSELQ SGQAQSKFKSEKQNRKQLELKVTSLEELT RVEKESELSKNLSSPEKKSAGRESQAEEEID RKSYGFEI.DKLRQLI.KKTRVSTDQASAVTSQ SGQAQSKFKSEKQNRKQLELKVTSLEELT RVEKESELSKNLSSPEKKSAGRESQAEEEID RKSYGFEI.DKLRQLI.KKTRVSTDQASALSELQ SCQAQSKFKSEKQNRKQLELKVTSLEELT RVEKESELSKNLSSPEKKSAGRESQAEEEID RKSYGFEI.DKLRQLI.KKTRVSTDQASALSELQ SCQAQSKFKSEKQNRKQLELKVTSLEELT RVEKESELSKNLSSPERSALSESDDFI RKSYGFEI.DKLRQLI.KKTRVSTDQASASASGQPQAPLNRE ESPRYPSGQVCARACEHLLASAGABAHLQQ EVCAQRDAYQQKI.VQLQEKSVCFACLALQ GTAALTKONEGEDESSEE EKAEERPRRPSQGQSASASSQOPQAPLNRE ESSTEAGSTVAGAALRSHHSQRSSLSGD DELFRGATLKALRRYAQPEEEDEVSNK			1	Į.	1		
NLQVLNLSYNLLGELYSSNYTGJEKVAL  NNHALIGPOTKELERLOTLOLRONALT  FPISIPDIFLSGNKL/TLPKINLTANLHLIS  LENLDILYFLIAUPHI, QILINQNEFSSCSG  FPISIPDIFLSGNKL/TLPKINLTANLHLIS  LENLDILYFLIAUPHI, QILINQNEFSSCSG  FPISIPSIE, GLEI-GJENMLQLAWETELCWD  GEISHL, QVLYLINHNYLNSI, POWYPSHI, TAI  GLSLNSNRLTVLSHNDLPANLEILDISRNQ)  APPPDYFVSLSVLDTHNKEFCELSTFIN  NHTINVTLAGPPADIYCVYPDSLSGVSLFCELSTFIN  NHTINVTLAGPPADIYCVYPDSLSGVSLFCELSTFIN  NHTINVTLAGPPADIYCVYPDSLSGVSLFCELSTFIN  NHTINVTLAGPPADIYCVYPDSLSGVSLFCELSTFIN  NHTINVTLAGPPADIYCVYPDSLSGVSLFCELSTFIN  NHTINVTLAGPPADIYCVYPDSLSGVSLFCELSTFIN  NHTINVTLAGPPADIYCVYPDSLSGVSLFCELSTFIN  NHTINVTLAGPPADIYCVYPDSLSGVSLFCELSTFIN  NHTINVTLAGPPADIYCVYPDSLSGVSLFCELSTFIN  NHTINVTLAGPPADIYCVYPDSLSGVSLFCELSTFIN  KYRGCGFICYTALQVSLVSKYBLYCHLOTY  NKLUCLYSBHTLAGWCLWCKKKEKER  KDNNIPLOTVALTS  GRANLASULGTHALSOQULKKEKER  KDNNIPLOTVALTS  GRANLASULGTHALSOQULKKEKER  KDNNIPLOTVALTS  GRANLASULGTHALSOQULKKEKER  KDNNIPLOTVALTS  GRANLASULGTHALSOQULKKEKER  KORGGTAATGNOAITEKTAPATMSIPTILV  AVBAYNYNGGYYKVGGKFGAAVLCHNIP  FREILLTISQQQYPVTVARIBYNSELMANT  STPYDDQRONWSIMFISERCAAVEFNKOVC  KCKNSTSLDAVISCHSCHAMPCAKGORKLLIVP  CAYGSEGVIGWTQATDGSLVFELMVRIN  STPYDDQRONWSIMFISERCAAVEFNKOVC  KCKNSTSLDAVISCHSCHAMPCAKGORKLLIVP  CAYGSEGVIGWTQATDGSLVFELMVRIN  STPYDDGRONWSIMFISERCAAVEFNKOVC  KCKNSTSLDAVISCHSCHAMPCAKGORKLLIVP  CAYGSEGVIGWTQATDGSLVFELMVRIN  STPYDDGRONWSIMFISERCAAVEFNKOVC  KCKNSTSLDAVISCHSCHAMPCAKGORKLLIVP  CAYGSEGVIGWTQATDGSLVFELMVRIN  STPYDDGRONWSIMFISERCAAVEFNKOVC  KCKNSTSLDAVISCHSCHAMPCAKGORKLLIVP  CAYGSEGVIGWTQATDGSLVFELMOVRIN  STPYDDGRONWSIMFISERCAAVEFNKOVC  KCKNSTSLDAVISCHSCHAMPCAKGORKLLIVP  CAYGAYGROYMSIMFISERCAAVEFNKOVC  KKNGSSDGHIVSSRIDSAAAVEFNKONSCLEDAVISCHSCHAMPCAKGHLLANGA  VSPPTSIFFKSGEBALRTISNLSSCLANTIS  AVKAKLISRMAKMGOPMCJUPPLOCONOLIN  AVTGULAGARONALPSTALSCNAMPCAKGHLLANGARONALPSTALSCHAMPCAKGHLLANGARONALPSTALSCLANGARONALPSTALSCHAMPCAKGHLLANGARONALPSTALSCHAMPCAKGHLLANGARONALPSTALSCHAMPCAKGHLLANGARONALPSTALSCHAMPCAKGHLANGARONALPSTALSCHAMPCAKGHLANGARONALPSTALSCHAMPCAKGHLANGARONALPSTALSCHAMPCAKGHLANGARONALPSTALSCHAMPCAKGHLANGAR			<del> </del>	-	sequence		
QKNHIAIIQDQTFKFLERLQTLIDARDNALT FIPSIPDIFLSGKILVTLYRKILQHLIINQNIFSSEG GTPSEPSIE.GLI-GIBENMOLA WETELCWD EGI.SH.QVI.VINHNYI.NSLPPGVPSHLTAI GLI-SINSNILTVI.SSIND.PAN ELD ISINXNI APPDVFVSLSV.DITHNKFICECELSTIN METHNYITAGPPADIYCVYPDSLSGVSLFSL. GCDEFSYLKSILKFSLFYCTYTT.TI-ELBISTRNQ APPDVFVSLSV.DITHNKFICECELSTIN METHNYITAGPPADIYCVYPDSLSGVSLFSL. GCDEFSYLKSILKFSLFYCTYTT.TI-ELBISTRNQ APPDVFVSLSV.DITHNKFICECELSTIN METHNYITAGPPADIYCVYPDSLSGVSLFSL. GCDEFSYLKSILKFSLFYCTYTT.TI-ELBISTRNQ APPDVFVSLSV.DITHNKFICECELSTIN METHNYITAGPPADIYCVYPDSLSGVSLFSL. GCDEFSYLKSILKFSLFYCTYTT.TI-ELBISTRNQ APPDVFVSLSV.DITHNKFICECELSTIN METHNYITAGPADIYCVYBLKISTRGYPQ QNEPNLCFEBDFVPQENRFANIQDAIPC KYDAYLCTSSKDFTWVQNALLKHLDTQY QNEPNLCFEBDFVPQENRFANIQSIRGFYQ KYDAYLCTSSKDFTWVQNALLKHLDTQY QNEPNLCFEBDFVPQENRFANIQSIRGFYQ YNEPNEDDQUDVGWFLKIKSQGILKKEKE KDNNIPLCTYATIS  816  2166 A 6646 I 3811 RDRAGGYRFAGKQHAAAAFYDVGGDRPW GNTQLFRRNPYKANAMFGAGDEDDTDTS KKQGQTAATGNQATFKTAPATMSTPTIL AVHAYRYTINGQVYVGQKGAAVLGHIT EYRILLYISQQQPVITVARHIVNFELLWRWIN STFYDDQRQNWSIMFESSKAAFYBNACHAUPL VAYAYAYLGYNGVYVGQKGAAVLGHIT EYRILLYISQQGPVITAGNAAFYBNACHAUPL CANGSEGYIGWTQATDSIL.VPGLADSA VSPTSIPKSGEPALRTINSNISLSQLAINTS AVKAKLISRMAKMGQPM.PILPPQLDSSD VAYTGWGGOPVYTTSVGYSLOPAHPAL MTSQAPOPSVTGLQAFSAALMQVSSLDSA VSPTSIPKSGEPALRTINSNISLSQLAINTS AVKAKLISRMAKMGQPM.PILPPQLOSD EDVNTLQGGGQVVTTSVGYSLOPAHPAL MTSQAPOPSVTGLQAFSAALMQVSSLDSA VSPTSIPKSGEPALRTINSNISHQRIUGE LKOGELEKSNREEDONIXGIELERNORYYS NLMMEKRNNSLOTATENTQARVHACKET QHOTTERNAVSVADKMHLMTKVEFLQ SAGNSMLPSNSVTMETSMINSNINGRUP LKOGELEKSNREEDONIXGIELERNORYYS NLMMEKRNNSLOTATENTQARVHACKET QHQLTESLKETULRGQLTKVQAKLSHLQ SEQAQSKTSEKGONRKOGLERNORYYS NLMMEKRNNSLOTATENTQARVHACKET QHQLTESLKETULRGQLTKVQAKLSHLQ SEQAQSKTSEKGONRKOGLERNORYYS LAMMEKRNNSLOTATENTQARVHACKET QHQLTESLKETULRGQLTKVGAKLSHLQ SEQAQSKTSEKGONRKOGLERNORYYS NLMMEKRNNSLOTATENTQARVHACKET QHQLTESLKETULRGQLTKVGAKLSHLG SEQAQSKTYCSKQNRKOGLERNORYSEEL SAGNSHARRAGGEDDDYSKKER ESSELAKERRPRPSOGQSASASSGQPQAPLNEE ESSPMYSEQVYCEAVPROQECEESISSEE EKAEERPRRPSOGQSASASSGQPQAPLNEE ESSPMYSEQVYCEAVPROGGEBEDEDEVSNK	ł	}	l	1	}		
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### STEPPLQDVGWFLIKLSQQILKKEKER KDNNIPLQTVATIS  ### STEPPLQDVG ### STEPPLQDVG ### STEPPLQDQAAAAFYDVGGDRPWI GNTQLPPRNPVKANAAMPGAGDEDTDTLS GGARLASLFGLDQAAAAGHGNEFFQYTAPP KKGQGTAATGNQATPKTAPATMSTPTILV AVHAYRYTINGQVPVQGKFGAAVLGNHTT EYRILLYISQQQPVTVARHIVNFELMVRPN STFYDDQRQNWSIMPESEKAAVEFNKQVC KCNSTSSLDAVLSQDLIVADGPAVEVGDSI VAYTGWLFQNHVLGQVFDSTANKDKLLR LGSGKVIKGWEDMLGMKKGGKRLLIVPI CAVGSEGVIGWTQATDSILVFEVEVRVKI KDSGSDGHSVSSRDSSAAPSPIPGADNLSAD VSPPTSIPFKSGEPALRTKSNSLSEQLAINTS AVKAKLISRMAKMGQPMLPILPPQLDSND EDVNTLQGGGQPVTPSVQPSLOPAHPALL MTSQAPQPSVTGLQAPSAALMQVSSLDSH VSGNAQSFQPYAGMQAYAYPQASAVTSQI PVRRLYPAPLSQPPHPQGSGDMASFLMITEZ QHNTERMAVSKVADKMDHLMTKVELQ SAGNSMLPSMYTWETSMIRNIQRIQEE LKQEILEKSNRIEEQNDKISELIERNQRYVE NLMMEKRNNSLQTATENTQARVLHAEQE KVTBELAAATAQVSHLQLKMTAHQKKETI QWQLTESLKETDLLRGQLTKVQAKLSELQ SEQAQSKFKSEKNKSQNKQLELKVTSLEEELI RVEKESLEKNLSERKKKSAQERSQAEEED RKSYQEELDKLRQLIKKTRVSTDQAAAEQ LVQAAELQTQWEAKCEHLLASAKDEHLQQ EVCAQRDAYQKLVULOKKVTSLEEELI RVEKESLEKNLSERKKKSAQERSQAEEED RKSYQEELDKLRQLIKKTRVSTDQAAAEQ LUTALTKQNEGNIKQLIEKVTSLEEELI RVEKESLEKNLSERKKKSAQERSQAEEED GEVCAQRDAYQKLVULOKSVCFACLAL QITALTKQNEGNIKQLIEKVTSLEEELI RVEKESLEKNLSERKKKSAQERSQAEEED BCATTIMNTIKMYTLOLLNQQEGEEESSSEE EKAEERPRRPSQEQSASASSGQPQAPLNRE ESPMVPSEQVVEEAVPLIPQALTTISQDGHE KGDSEAEALSIERDGSLPPELSCIPSHRVLG TSIPPEPLGPVSMDSECEESLAASPMAAKP PSGKKVCVREVAPDGPLQEESTSRLS.TSOMP PGGRKVCVREVAPDGPLQEESTRSLSTSOMP GDPLALGPESPGEPQPPQLKKDDVTSSTGP ELSSTEAGSTVAGAALRFSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK		1	1	1			
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S16   2166   A   6646   I   3811   RDRAGVRPAGKQHAAAAFYDVGGDRPW GNTQLPPRNPVKANAMFGAGDEDDTDLS GGARLASLFGLDQAAAGHGNEFFQYTAFK KKGQGTAATGNQATPKTAPATMSTPTILV AVHAYRYTINGQYVKQGKFGAAVLGNHTT EYRILLYISQQQPVTVARHIVNFELMWRPN STFYDDQRQNWSIMFESEKAAVEFNKQVC KCNSTSSLDAVLSQDLIVADGPAVEVGDSI VAYTGWLFQNHVLGQVFDSTANKDKLLR LGSGKVIKGWEGMILGMKKGGKRLLVPP CAVGSEGVIGWTQATDSILVFEVEVRVK KDSGSDGHSVSRSAAPSPIPGADNLSAAD VSPPTSIPFKSGEPALRTKSNSLSEQLAINTS AVKAKLISRMAKMGQPMLPILLPPQLDSND EDVNTLQGGGQPVVTPSVQPSLQPAHFALL MTSQAPQPSVTGLQAPSAALMQVSSLDSH VSGNAQSFQPYAMFALM QHNTEIRMAVSKVADKMDHLMTKVEELQ SAGNSMLPSMYTMETSMIMSNIQRIIQER LKQEILEKSNRIEEQNDKISELIERNQRYVE NLMMEKRNNSLYGTATENTOARVHAEQE KVTEELAAATAQVSHLQLKMTRYSTDQAAASE KVTEELAAATAQVSHLQLKMTAVSTDQAASCKSTL RVEKESLEKNLSERKKKSAQERSQAEEED RKSYQEELDKLQLKKTRYSTDQAAASE LVQAELQTQWEAKCEHLLASAKDEHLQQ EVCAQRDAYQKLVQLCEKSVGFACLAL QITALTKQNSCHLKKTRYSTDQAASC LVQAELQTQWEAKCEHLLASAKDEHLQQ EVCAQRDAYQKLVQLCEKSVGFACLAL QITALTKQNSCHLKKTRYSTDQAASC LVQAELQTQWEAKCEHLLASAKDEHLQQ EVCAQRDAYQKLVQLCEKSVGFACLAL QITALTKQNSCHLKKERVSTDQAAASC LVQAELQTQWEAKCEHLLASAKDEHLQQ EVCAQRDAYQKLVQLCEKSVGFACLAL GTIMNTIKMVTLQLLQLQCEGECESSSEE EKAEERPRRPSQEQSASASSGQPQAPLNRE ESPMVPSEQVVEEAVPLPQALTTSQDGHR KGDSEAEALSERDGSLPPELSCFSHRVLG TSIPPEPLGPVSMDSECEELSAASPMAAKP PSGKKVCVREVAPDGPLQESSTRLS.TTSUPP GDFLALGPESPGEPQPPQLKKDDVTSSTGP ELSSTEAGSTVAGAALRFKAQPEEEDEDEVSMK							
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GGARLASLFGLDQAAAGHGNEFFQYTAPK KKGQGTAATGNQATPKTAPATMSTPTILV AVHAYRYTNGQYVKQGKFGAAVLGNHTI EYRILLYISQQQPVTVARIHVNFELMVRPN STFYDDQRQNSIMFESSEKAVEFNKQVC KCNSTSSLDAVLSQDLIVADGPAVEVGDSI VAYTGWLFQNHVLGQVFDSTANKDKLLR LGSGKVIKGWEDGMLGMKKGGKRLIVPI CAVGSEGVIGWTQATDSILVFEVEVRVKI KDSGSDGHSVSSRDSAAPSPIPGADNLSAD VSPPTSIPFKSGEPALRIKSNSLSEQLAINTS AVKAKLISRMAKMGQPMLPILPPQLDSND EDVNTLQGGOPVVTPSVOPSLQPAHPALI MTSQAPQPSVTGLQAPSAALMQVSSLDSH VSGNAQSFQPYAGMQAYAYPQASAVTSQI PVRPLYPAPLSOPPHFQGSDMASFLMTEA QHNTEERMAVSKVADKMDHLMTKVEELQ SAGNSMLIPSMSVTMETSMIMSNIQRIIQEB LKQEILEKSNRIEEQNDKISELIERNQRYVE NLMMEKRNNSLQTATENTQARVLHAEQE KVTEELAAATQVSHLQLKMTAHQKKETI QMQLTESLKETDLLRGQLTKVQAKLSELQ SEQAQSKFKSEKQNRKQLELKVTSLEEELI RVEKRSLEKKNLSERKKRSAQERSQAEEGEID RKSYQEELDKLRQLLKKTRVSTDQAAAEQ LVQAELQTQWEAKCHLLASAKDEHLQQ EVCAQRDAYQQKLVQLQEKSVCFALCLAL QITALTKQNEQHIKELEKNRSQMSGVEAA. DPSEKVKKIMNQVFQSLREFELEESYNGF LGTIMNTIKMVTLQLLNQEQEKEESSSEE EKAEERPRRPSQEQSASASSGQPQAPLNRE ESPMYPSEQVVEEAVPLPQAALTTSQDGHF KGDSEAEALSEIKDGSLPPELSCIPSHRVLG TSIPPEPLGPVSMDSECESLAASPMAAKPP PSGKVCVYREVAPDGPLQESSTRLSLTSVDP GDPLALGPESPGEPPPQLKKDDVTSSTGP ELSSTEAGSTVAGAAARPSSHSQRSSLSGD DELFKGATLKLARPKAQEEEEDEDEVSMK	810	2100	A	0040	1	3811	
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RKSYQEELDKLRQLLKKTRVSTDQAAAEQ LVQAELQTQWEAKCEHLLASAKDEHLQQ' EVCAQRDAYQQKLVQLQEKSVCFA\CLAL QITALTKQNEQHIKELEKNKSQMSGVEAA. DPSEKVKKIMNQVFQSLRREFELEESYNGR LGTIMNTIKMVTLQLLNQQEQEKEESSSEE EKAERPRRPSQEQSASASSGQPQAPLNRE ESPMVPSEQVVEEAVPLPPQALTTSQDGHR KGDSEAEALSEIKDGSLPPELSCIPSHRVLG TSIPPEPLGPVSMDSECEESLAASPMAAK\P PSGK\VCVREVAPDGPLQESSTRLSLTS\DPI GDPLALGPESPGEPQPPQLKKDDVTSSTGP ELSSTEAGSTVAGAALRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK							RVEKESLEKNLSERKKKSAQERSQAEEEIDEI
LVQAELQTQWEAKCEHLLASAKDEHLQQ EVCAQRDAYQQKLVQLQEKSVCFA\CLAL QITALTKQNEQHIKELEKNKSQMSGVEAA. DPSEKVKKIMNQVFQSLRREFELEESYNGR LGTIMNTIKMVTLQLLNQQEQEKEESSSEE EKAERPRRPSQEQSASASSGQPQAPLNRE ESPMVPSEQVVEEAVPLPPQALTTSQDGHR KGDSEAEALSEIKDGSLPPELSCIPSHRVLG TSIPPEPLGPVSMDSECEESLAASPMAAK\P PSGK\VCVREVAPDGPLQESSTRLSLTS\DPI GDPLALGPESPGEPQPPQLKKDDVTSSTGP ELSSTEAGSTVAGAALRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK			1			:	
EVCAQRDAYQQKLVQLQEKSVCFA\CLAL QITALTKQNEQHIKELEKNKSQMSGVEAA, DPSEKVKKIMNQVFQSLRREFELEESYNGR LGTIMNTIKMVTLQLLNQQEQEKEESSSEE EKAEERPRRPSQEQSASASSGQPQAPLNRE ESPMVPSEQVVEEAVPLPPQALTTSQDGHR KGDSEAELSEIKDGSLPPELSCIPSHRVLG TSIPPEPLGPVSMDSECEESLAASPMAAK\P. PSGK\VCVREVAPDGPLQESSTRLSLTS\DPI GDPLALGPESPGEPQPPQLKKDDVTSSTGP ELSSTEAGSTVAGAALRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK			1			'	
QITALTKQNEQHIKELEKNKSQMSGVEAA.  DPSEKVKKIMNQVFQSLRREFELEESYNGR LGTIMNTIKMVTLQLLNQQEQEKEESSSEE EKAEERPRRPSQEQSASASSGQPQAPLNRE ESPMVPSEQVVEEAVPLPPQALTTSQDGHR KGDSEAEALSEIKDGSLPPELSCIPSHRVLG TSIPPEPLGPVSMDSECEESLAASPMAAK\P PSGK\VCVREVAPDGPLQESSTRLSLTS\DPI GDPLALGPESPGEPQPPQLKKDDVTSSTGP ELSSTEAGSTVAGAALRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK				1			
DPSEKVKKIMNQVFQSLRREFELEESYNGR LGTIMNTIKMVTLQLLNQQEQEKEESSSEE EKAEERPRRPSQEQSASASSGQPQAPLNRE ESPMVPSEQVVEEAVPLPPQALTTSQDGHR KGDSEAEALSEIKDGSLPPELSCIPSHRVLG TSIPPEPLGPVSMDSECEESLAASPMAAK\P. PSGK\VCVREVAPDGPLQESSTRLSLTS\DPH GDPLALGPESPGEPQPPQLKKDDVTSSTGP ELSSTEAGSTVAGAALRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK							
LGTIMNTIKMVTLQLLNQQEQEKEESSSEE EKAEERPRRPSQEQSASASSGQPQAPLNRE ESPMVPSEQVVEEAVPLPPQALTTSQDGHR KGDSEAEALSEIKDGSLPPELSCIPSHRVLG TSIPPEPLGPVSMDSECEESLAASPMAAK\P. PSGK\VCVREVAPDGPLQESSTRLSLTS\DPI GDPLALGPESPGEPQPPQLKKDDVTSSTGP. ELSSTEAGSTVAGAALRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK							
EKAEERPRRPSQEQSASASSGQPQAPLNRE ESPMVPSEQVVEEAVPLPPQALTTSQDGHR KGDSEAEALSEIKDGSLPPELSCIPSHRVLG TSIPPEPLGPVSMDSECEESLAASPMAAK\P. PSGK\VCVREVAPDGPLQESSTRLSLTS\DPI GDPLALGPESPGEPQPPQLKKDDVTSSTGP. ELSSTEAGSTVAGAALRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK				}	]		Droew v Kanving v ryslakkerelees y NGRTI
ESPMVPSEQVVEEAVPLPPQALTTSQDGHR KGDSEAEALSEIKDGSLPPELSCIPSHRVLG TSIPPEPLGPVSMDSECEESLAASPMAAK\P. PSGK\VCVREVAPDGPLQESSTRLSLTS\DPI GDPLALGPESPGEPQPPQLKKDDVTSSTGP: ELSSTEAGSTVAGAALRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK							LGIIMNIIKMVILQLLNQQEQEKEESSSEEEE
KGDSEAEALSEIKDGSLPPELSCIPSHRVLG TSIPPEPLGPVSMDSECEESLAASPMAAK\P. PSGK\VCVREVAPDGPLQESSTRLSLTS\DPI GDPLALGPESPGEPQPPQLKKDDVTSSTGP ELSSTEAGSTVAGAALRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK			ĺ				
TSIPPEPLGPVSMDSECEESLAASPMAAK\P. PSGK\VCVREVAPDGPLQESSTRLSLTS\DPI GDPLALGPESPGEPQPPQLKKDDVTSSTGP. ELSSTEAGSTVAGAALRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK							ESPMVPSEQVVEEAVPLPPQALTTSQDGHRR
PSGK\VCVREVAPDGPLQESSTRLSLTS\DPI GDPLALGPESPGEPQPPQLKKDDVTSSTGP ELSSTEAGSTVAGAALRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK							KGDSEAEALSEIKDGSLPPELSCIPSHRVLGPP
PSGK\VCVREVAPDGPLQESSTRLSLTS\DPI GDPLALGPESPGEPQPPQLKKDDVTSSTGP ELSSTEAGSTVAGAALRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK			l	J I			TSIPPEPLGPVSMDSECEESLAASPMAAK\PDN
GDPLALGPESPGEPQPPQLKKDDVTSSTGP ELSSTEAGSTVAGAALRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK							
ELSSTEAGSTVAGAÅLRPSHHSQRSSLSGD DELFKGATLKALRPKAQPEEEDEDEVSMK				[	,		
DELFKGATLKALRPKAQPEEEDEDEVSMK				<u>,</u>			
				j j	:		
PPPTPLFGDDDDDDDDDDDDDDDDDDDDDDDDDDDDDDDDDDDD	015	01.55		6640		1070	
817 2167 A 6649 63 1073 FFRSSSDNGSPIRQYE/HSTPAHQGPVMGLE	81./	2167	A	6649	6.5	1073	FFRSSSDNGSPIRQYE/HSTPAHQGPVMGLEG

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  KS/ARNSQLRIVLVGKTGAGKSATGNSILGRK VFHSGTAAKSITKKCEKRSSSWKETELVVVD TPGIFDTEVPNAETSKEIIRCILLTSPGPHALLL VVPLGRYTEEEHKATEKILKMFGERARSFMIL IFTRKDDLGDTNLHDYLREAPEDIQDLMDIFG DRYCALNNKATGAEQEAQRAQLLGLIQRVV RENKEGCYTNRMYQRAEEEIQKQTQAMQEL HRVELEREKARIREEYEEKIRKLEDKVEQEKR KKQMEKKLAEQEAHYAVRQQRARTEVESKD GILELIMTALQIASFILLRLFAED
818	2168	A	6660	357	1890	APSGSWTRVVLTLDPCSLRSRSPRSLLDPGMP GISARGLSHEGRKQLAVNLTRVLALYRSILDA YIIEFF'TDNLWDTLPCSWQEALDGLKPPQLA TMLLGMPGEGEVVRYRSVWPLTLLALKSTA CALAFTRMPGFQTPSEFLENPSQSSRLTAPFR KHVRPKKQHEIRRLGELVKKLSDFT/GLHPGC RRGLRPG'HLSRFMALGLGLMVKSIEGDQRL VERAQRLDQELLQALEKEEKRNPQVVQTSPR HSPHHVVRWVDPTALCEELLLPLENPCQGRA RLLLTGLHACG'OLSVALLRHFSCCPEVVALA SVGCCYMKLSDPGGYPLSQWVAGLPGYELP YRLREGACHALEEYAERLQKAGPGLRTHCY RAALETVIRRARPELRRPGVQGIPRVHELKIEE YVQRGLQRVGLDPQLPLNLAALQAHLAQEN RVVAFFSLALLLAPLVETLILLDRLLYLQEQA LSP\GFHAELLPIFSPELSPRNLVLVATKMPLG QALSVLETEDS
819	2169	A	6661	65	2686	SGSGHCLAEAASMGPWGWKLRWTVALLLA AAGTAVGDRCERNEFQCQDGKCISYKWVCD GSAECQDGSDESQETCLSVTCKSGDFSCGGR VNRCIPQFWRCDGQVDCDNGSDEQGCPPKTC SQDEFRCHDGKCISRQFVCDSDRDCLDGSDE ASCPVLTCGPASFQCNSSTCIPQLWACDNDPD CEDGSDEWPQRCRGLYVFQGDSSPCSAFEFH CLSGECIHSSWRCDGGPDCKDKSDEENCAVA TCRPDEFQCSDGNCIHGSRQCDREYDCKDMS DEVGCVNVTLCEGPNKFKCHSGECITLDKVC NMARDCRDWSDEPIKECGTNECLDNNGGCS HVCNDLKIGYECLCPDGFQLVAQRRCEDIDE CQDPDTCSQLCVNLEGGYKCQCEEGFQLDPH TKACKAVGSIAYLFFTNRHEVRKMTLDRSEY TSLIPNLRNVVALDTEVASNRIYWSDLSQRMI CSTQLDRAHGVSSYDTVISRDIQAPDGLAVD WIHSNIYWTDSVLGTVSVADTKGVKRKTLFR ENGSKPRAIVVDPVHGFMYWTDWGTPAKIK KGGLNGVDIYSLVTENIQWPNGITLDLLSGRL YWVDSKLHSISSIDVNGGNRKTILEDEKRLAH PFSLAVFEDKVFWTDIINEAIFSANRLTGSDV NLLAENLLSPEDMVLFHNLTQPRGVNWCERT TLSNGGCQYLCLPAPQINPHSPKFTCACPDGM LLAR\DMRSCLTEG\EAAVATQETSTVRLKVS STAVRTQHTTTRPVPDTSRLPGATPGLTTVEI VTMSHQALGDVAG\RGN\EKKPSSVRALSIVL PIVLLVFLCLGVFLLWKNWRLKNINSINFDNP VYQKTTEDEVHICHNQDGYSYPSRQMVSLED DVA
820	2170	A	6666	17	4146	ERGISSQIKGMKSGSGGGSPTSLWGLLFLSAA LSLWPTSGEICGPGIDIRNDYQQLKRLENCTVI EGYLHILLISKAEDYRSYRFPKLTVITEYLLLF RVAGLESLGDLFPNLTVIRGWKLFYNYALVIF

SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence	ł	l	914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
		'		amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
		ļ	1	residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
	l	1	1	peptide	5 - 4	/=possible nucleotide deletion, \=possible
		İ	l .	sequence		nucleotide insertion
	<u> </u>	·	<del> </del>			EMTNLKDIGLYNLRNITRG\AIRIEKNADLCY
						STVDWSLILDAVSNNYIVGNKPPKECGDLCF
	ı	1	1		}	GTMEEKPMCEKTTINNEYNYRCWTTNRCQI
		1				MCPSTCGKRACTENNECCHPECLGSCSAPDI
						DTACVACRHYYYAGVCVPACPPNTYRFEGV
i	1	ł	ł		}	RCVDRDFCANILSAESSDSEGFVIHDGECMO
						CPSGFIRNGSQSMYCIPCEGPCPKVCEEEKKT
		1				KTIDSVTSAQMLQGCTIFKGNLLINIRRGNNI
	l	1			1	SELENFMGLIEVVTGYVKIRHSHALVSLSFLI
						NLRLILGEEQLEGNYSFYVLDNQNLQQLWD
	ĺ	ĺ	ĺ			WDHRNLTIKAGKMYFAFNPKLCVSEIYRME
						VTGTKGRQSKGDINTRNNGERASCESDVLHI
				Ì	ľ	TSTTTSKNRIIITWHRYRPPDYRDLISFTVYYI
	[					EAPFKNVTEYDGQDACGSNSWNMVDVDLP
						NKDVEPGILLHGLKPWTQYAVYVKAVTLTN
					ļ	VENDHIRGAKSEILYIRTNASVPSIPLDVLSAS
	ĺ		ĺ			NSSSQLIVKWNPPSLPNGNLSYYIVRWQRQP
		1		:		QDGYLYRHNYCSKDKIPIRKYADGTIDIEEV
:						ENPKTEVCGGEKGPCCACPKTEAEKQAEKE
						AEYRKVFENFLHNSIFVPRPERKRRDVMQVA
	ĺ	l			i	NTTMSSRSRNTTAADTYNITDPEELETEYPFF
			İ			ESRVDNKERTVISNLRPFTLYRIDIHSCNHEA
						KLGCSASNFVFARTMPAEGADDIPGPVTWEI
	ĺ	İ	ĺ	ĺ	1	RPENSIFLKWPEPENPNGLILMYEIKYGSQVE
						DQRECVSRQEYRKYGGAKLNRLNPGNYTA
						QATSLSGNGSWTDPVFFYVQAKRYENFIHLI
í	ì	ĺ	!		Ì	ALPVAVLLIVGGLVIMLYVFHRKRNNSRLGN
			l			GVLYASVNPEYFSAADVYVPDEWEVAREKI
						MSRELGQGSFGMVYEGVAKGVVKDEPETR
[		1	ĺ			AIKTVNEAASMRERIEFLNEASVMKEFNCHF
					1	VVRLLGVVSQGQPTLVIMELMTRGDLKSYL
			J			SLRPEMENNPVLAPPSLSKMIQMAGEIADGM
			[			AYLNANKFVHRDLAARNCMVAEDFTVKIGI
1						FGMTRDIYETDYYRKGGKGLLPVRWMSPES
			ł			KDGVFTTYSDVWSFGVVLWEIATLAEQPYQ
		1				GLSNEQVLRFV\MEGGLLDKPDNCPDMLFEI
1			1			MRMCWQYNPKMRPSFLEIISSIKEEMEPGFR
						VSFYYSEENKLPEPEELDLEPENMESVPLDPS
						ASSSSLPLPDRHSGHKAENGPGPGVLVLRAS
		1		1		DERQPYAHMNGGRKNERALPLPQSSTC
821	2171	Α	6691	106	825	GRVLFRGCGVGHKGQVLMGTFILAQDWLSE
	-					SNHVFCVSSMLRLOKRLASSVLRCGKKKVW
1		1	[	ľ		LDPNETNEIANANSRQQIRKLIKDGLIIRKPVI
						VHSRARCRKNTLARRKGRHMGIGKRKGTAN
						ARMPEKVTWMRRMRILRRLLRRYRES/KRY
Í			[			ESKKIDRHMYHSLYLKVKGNVFKNKRILME
						IHKLKADKARKKLLADQAEARRSKTKEARK
		}	]			RREERLQAKKEEIIKTLSKEEETKK
822	2172	A	6715	772	21	DFRPGLLLPRKKKMFGFHKPKMYRSIEGC\C
	4114	A	6,12	114	<del>-</del> 1	
1						SGAKSSSS\RFTDSKRYEK\DFQ\SCFGLHETR SGDI\CNA\CVLL\LKRWKKLPAGSKK\NWNH
r						
		I		1		VVDARAGPS\LKTTLKPKKVKTL\SGNRIK\ST
				1	l l	QISKLQKEFKR\HNSDAHSTTS\SASP\AQSPLF
						### 10 PD 11 m 0 0 D # 0 1 1
						TYWKRQKICCGI\IYKGRFGEVLIDTHLFKPC
						TYWKRQKICCGI\IYKGRFGEVLIDTHLFKPC SNKKA\AAEKPEEQGPEPLPISTQEWVTEVFM
823	2173	A	6727	3	4063	TVNQFRWTGSDTGVGFPGSNRNHPVFSFLDL TYWKRQKICCGI\IYKGRFGEVLIDTHLFKPCG SNKKA\AAEKPEEQGPEPLPISTQEWVTEVFM PYLATLQLDSSLLIPPKYQTPPAAAQGQATPC
823	2173	Ą	6727	3	4063	TYWKRQKICCGI\IYKGRFGEVLIDTHLFKPC SNKKA\AAEKPEEQGPEPLPISTQEWVTEVFM
823	2173	Ā	6727	3	4063	TYWKRQKICCGNYKGRFGEVLIDTHLFKPCG SNKKA\AAEKPEEQGPEPLPISTQEWVTEVFM PYLATLQLDSSLLIPPKYQTPPAAAQGQATPC

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  SSQPSQDGQESNVPSVGSLADPDYLNTPQMN TPVTLNSAAPASNSGAGVLPSPATPRFSVPTP RTPRTPRTPRTPRGGGTASGQGSVKYDSTDQGSP ASTPSTTRPLNSVEPATMQPIPEAHSLYVTLIL SDSVMNIFKDRNFDSCCICACNMNIKGADVG LYIPDSSNEDQYRCTCGFSAIMNRKLGYNSGL FLEDELDIFGKNSDIGQAAERRLMMCQSTFL PQVEGTKKPQEPPISLLLLLQNQHTQPFASLN FLDYISSNNRQTLPCVSWSYDRVQADNNDY WTECFNALEQGRQYVDNPTGGKVDEALVRS ATVHSWPHSNVLDISMLSSQDVVRMLLSLQP FLQDAIQKKRTGRTWENIQHVQGPLTWQQFH KMAGRGTYGSEESPEPLPIPTLLVGYDKDFLT ISPFSLPFWERLLLDPYGGHRDVAYIVVCPEN EALLEGAKTFFRDLSAVYEMCRLGQHKPICK VLRDGIMRVGKTVAQKLTDELVSEWFNQPW SGEENDNHSRLKLYAQVCRHHLAPYLATLQL DSSLLIPPKYQTPPAAAQGATPGNAGPLAPN GSAAPPAGSAFNPTSNSSSTNPAASSSASGSV PPVSSSASAPGISQISTTSSSGFSGSVGGQNPST GGISADRTQGNIGCGGDTDPGQSSSQPSQDG QESVTERERIGIPTEPDSADSHAHPPAVVIYM VDPFTYAAEEDSTSGNFWLLSLMRCYTEMLD NLPEHMRNSFILQIVPCQYMLQTMKDEQVFY IQYLKSMAFSVYCQCRRPLPTQIHIKSLTGFGP AASIEMTLKNPERPSPIQLYSPPFILAPIKDKQT ELGETFGEASQKYNVLFVGYCLSHDQRWLL ASCTDLHGELLETCVVNIALPNRSRRSKVSAR KIGLQKLWEWCIGIVQMTSLPWRVVIGRLGR LGHGELKDWSILLGECSLQTISKKLKDVCRM CGISAADSPSILSACLVAMEPQGSFVVMPDAV
						TMGSVFGRSTALNMQSSQLNTPQDASCTHIL VFPTSSTIQVAPANYPNEDGFSPNNDDMFVDL PFPDDMDNDIGILMTGNLHSSPNSSPVPSPGSP SGIGVGSHFQHSRSQGERLLSREAPEELKQQP LALGYFVSTAKAENLPQWFWSSCPQAQN\QC PLFLKASLHHHISVAQTDELLPARNSQRVPHP LDSKTTSDVLRFVLEQYNALSWLTCNPATQD RTSCLPVHFVVLTQLYNAIMNIL
824	2174	A	6732	2440	365	VEEGLGRRRTPPGGRRGPVTPARPGPDSVRR RLLPPSSAAAFSSHRHNILCSRRRGGGGGG GGGGGTIKRPGITGPTAATSPSGEPGNAASAP LSLLSPFPGQTTYQHPGVAEPSAYGGRDVAC ASLVFGRLQHRGGDRKRGLLGRSSGDAASD QPFRCRSGSTAGRLVKQMDFTEAYADTCSTV GLAAREGNVKVLRKLLKKGRSVDVADNRG WMPIHEAAYHNSVECLQMLINADSSENYIKM KTFEGFCALHLAASQGHWKIVQILLEAGADP NATTLEETTPLFLAVENGQIDVLRLLLQHGAN VNGSHSMCGWNSLHQASFQENAEIIKLLLRK GANKECQDDFGITPLFVAAQYGKLESL\SILIS SG\ANVNCQALDKATPLFIAAQEGHTKCVELL LSSGADPDLYCNEDSWQLPIHAAAQMGHTKI LDLLIPLTNRACDTGLNKVSPVYSAVFGGHE DCLEILLRNGYSPDAQACLVFGFSSPVCMAFQ KDCEFFGIVNILLKYGAQINELHLAYCLKYEK FSIFRYFLRKGCSLGPWNHIYEFVNHAIKAQA KYKEWLPHLLVAGFDPLILLCNSWIDSVSIDT LIFTLEFTNWKTLAPAVERMLSARASNAWIL QQHIATVPSLTHLCRLEIRSSLKSERLRSDSYIS

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825	2175	A	6735	277	1252	G RIMGLFDRGVQMLLTTVGAFAAFSLMTIAVG TDYWLYSRGVCKTKSVSENETSKKNEEVMT HSGLWRTCCLEGNFKGLCKQIDHFPEDADYE ADTAEYFLRAVRASSIFPILSVILLFMGGLCIA ASEFYKTRHNIILSAGIFFVSAGLSNIIGIIVYIS ANAGDPSKSDSKKNSYSYGWSFYFGALSFIIA EMVGVLAVHMFIDRHKQLRATARA\TDYLQ ASAITRIPSYRYRYQRRSRSSRSTEPSHSRDA SPVGIKGFNTLPSTEISMYTLSRDPLKAATTPT ATYNSDRDNSFLQVHNCIQKENKDSLHSNTA NRRTTPV
826	2176	A	6744	3	5177	SDDLRTGLFQDVQDAESLKLPGVYEVLFYNE TEDCPGMMLWRYPEPRGLTLVRITPVPFNTT EDPDISTADLGDVLQDPCSLEYWDELQKVFV AFREFNLSESKVCELQLPDINLVNDQKKLVSS DLWRIVLNSSQNGADDQSSASESGSTCDPL VTPTALAACTRVDSCFTPWFVPSLCVSFQFAH LEFHLCHHLDQLGTAAPQYLQPFVSDRNMPS ELEYMIVSFREPHMYLRQWNNGSVCQEIQFL AQADCKLLECRNVTMQSVVKPFSIFGQMAVS SDVVEKLLDCTVIVDSVFVNLGQHVVHSLNT AIQAWQQNKCPEVEELVFSHFVICNDTQETL RFGQVDTDENILLASLHSHQYSWRSHKSPQL LHICIEGWGNWRWSEPFSVDHAGTFIRTIQYR GRTASLIIKVQQLNGVQKQIIICGRQIICSYLSQ SIELKVVQHYIGQDGQAVVREHFDCLTAKQK LPSYILENNELTELCVKAKGDEDWSRDVCLE SKAPEYSIVIQVPSSNSSIIYVWCTVLTLEPNS QVQQRMIVFSPLFIMRSHLPDPIIIHLEKRSLGL SETQIIPGKGQEKPLQNIEPDLVHHLTFQAREE YDPSDCAVPISTSLIKQIATKVHPGGTVNQILD EFYGPEKSLQPIWPYNKKDSDRNEQLSQWDS PMRVKLSIWKPYVRTLLIELLPWALLINESKW DLWLFEGEKIVLQVPAGKIIIPPNFQEAFQIGIY WANTNTVHKSVAIKLVHNLTSPKWKDGGNG EVVTLDEEAFVDTEIRLGAFPGHQKLCQFCIS SMVQQGIQIIQIEDKTTIINNTPYQIFYKPQLSV CNPHSGKEYFRVPDSATFSICPGGEQPAMKSS SLPCWDLMPDISQSVLDASLLQKQIMLGFSPA PGADSSQCWSLPAIVRPEFPRQSVAVPLGNFR ENGFCTRAIVLTYQEHLGVTYLTLSEDPSPRV IIHNRCPVKMLIKENIKDIPKFEVYCKKIPSECS IHHELYHQISSYPDCKTKDLLPSLLLRVEPLDE VTTEWSDAIDINSQGTQVVFLTGFGYVYVDV VHQCGTVFITVAPEGKAGPILTNTNRAPEKIV TF/KMFITQLSLAVFDDLTHHKASAELLRLTL DNIFLCVAPGAGPLPGEEPVAALFELYCVEIC CGDLQLDNQLYNKSNFHFAVLVCQGEKAEPI QCSKMQSLLISNKELEEYKEKCFIKLCITLNEG KSILCDINEFSFELKPARLYVEDTFVYYIKTLF DTYLPNSRLAGHSTHLSGGKQVLPMQVTQH ARALVNPVKLRKLVIQPVNLLVSIHASLKLYI ASDHTPLSFSVFERGPIFTTARQLVHALAMMHY AAGALFRAGWVVGSLDILGSPASLVRSIGNG VADFFRLPYEGLTRGPGAFVSGVSRGTTSFVK HISKGTLTSITNLATSLARNMDRLSLDEEHYN RQEEWRRQLPESLGEGLRQGLSRLGISLLGAI AGIVDQPMQNFQKTSEAQASAGHKAKGVISG

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  VGKGIMGVFTKPIGGAAELVSQTGYGILHGA GLSQLPKQRHQPSDVHADQAPNSHVKYVW KMLQSLGRPEVHMALDVVLVRGSGQEHEGC LLLTSEVLFVVSVEDTQQAFPVTEIDCAQD
827	2177	A	6748	2	1662	SKQNNLLTVQLKQPRVACDVEVDGVRERLSE QQYNRLVDYITKTSCHLAPSCSSMQIPCPVVA AEPPPSTVKTYHYLVDPHFAQVFLSKFTMVK NKALRKGFP FVGAPRRGNPFGSPGNPGRHQGPCHRPRGTK
						ASGVSPTLWRPQAAATGLEMPSSGRALLDSP LDSGSLTSLDSSVFCSEGEGEPLALGDCFTVN VGGSRFVLSQQALSCFPHTRLGKLAVVVASY RRPGALAAVPSPLELCDDANPVDNEYFFDRS SQAFRYVLHYYRTGRLHVMEQLCALSFLQEI QYWGIDELSIDSCCRDRYFRRKELSETLDFKK DTEDQESQHESEQDFSQGPCPTVRQKLWNIL EKPGSSTAARIFGVISIIFVGVSIINMALMSAEL SWLDLQLLEILEYVCISWFTGEFVLRFLCVRD RCRFLRKVPNIIDLLAILPFYITLLVESLSG\SQT TQEL\'ENVGAHCPGCLRLLRAL\RMLKAWGR HSTGLRSLGMTITQCYEEVGLLLLFLSVGISIF STVEYFAEQSIPDTTFTSVPCAWWWATTSMT TVGYGDIRPDTTTGKIVAFMCILSGILVLALPI AIINDRFSACYFTLKLKEAAVRQREALKKLTK NIATDSYISVNLRDVYARSIMEMLRLKGRER ASTRSSGGDDFWF
828	2178	A	6786	5672	1360	GTHPASSGPVPLPPAAVSAATREELGEPVFFV TASSGFQSMHSSNPKVRSSPSGNTQSSPKSKQ EVMVRPPTVMSPSGNPQLDSKFSNQGKQGGS ASQSQPSPCDSKSGGHTPKALPGPGGSMGLK NGAGNGAKGKGKRERSISADSFDQRDPGTPN DDSDIKECNSADHIKSQDSQHTPHSMTPSNAT APRSSTPPHGQTTATEPTPAQKTPAKVVYVFS TEMANKAAEAVLKGQVETIVSFHIQNISNNK TERSTAPLNTQISALRNDPKPLPQQPPAPANQ DQNSSQNTRLQPTPPIPAPAPKPAAPPRPLDRE SPGVENKLIPSVGSPASSTPLPPDGTGPNSTPN NRAVTPVSQGSNSSSADPKAPPPPPVSSGEPPT LGENPDGLSQEQLEHRERSLQTLRDIQRMLFP DEKEPTGAQSGGPQQNPGVLDGPQKKPEGPI QAMMAQSQSLGKGPGPRTDVGAPFGPQGHR DVPFSPDEMVPPSMNSQSGTIGPDHLDHMTP EQIAWLKLQQEFYEEKRRKPEQVVVQQCSLQ DMMVHQHGPRGVVRGPPPPYQMTPSEGWAP GGTEPFSDGINMPHSLPPRGMAPHPNMPGSQ MRLPGFAGMINSEMEGPNVPNPASRPGLSGV SWPDDVPKIPDGRNFPPGQGIFSGPGRGERFP NPQGLSEEMFQQQLAEKQLGLPPGMAMEGIR PSMEMNRMIPGSQRHMEPGNNPIFPRIPVEGP LSPSRGDFPKGIPPQMGPGRELEFGMVPSGM KGDVNLNVNMGSNSQMIPQKMREAGAGPEE MLKLRPGGSDMLPAQQKMVPLPFGEHPQQE YGMGPRPFLPMSQGPGSNSGLRNLREPIGPDQ RTNSRLSHMPPLPLNPSSNPTSLNTAPPVQRG LGRKPLDISVAGSQVHSPGINPLKSPTMHQVQ SPMLGSPSGNLKSPQTPSQLAGMLAGPAAAA SIKSPPVLGSAAASPVHLKSPSLPAPSPGWTSS PEPPLQSPGIPPNHKAPLTMASPAMLGNVESG GPPPPTASQPASVNIPGSSLPSSTPYTMPPEPTL SQNPLSIMMSRVMSKFAMPS\SNPGYNHDAI

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  KTVASSDDDSPPARSPNLPSMNNMPGMGINT QNPRISGPNPVVPMPTLSPMGMTQPLSHSNQ MPSPNAVGPNIPPHGVPMGPGLMSHNPIMGH GSQEPPMVPQGRMGFPQGFPPVQSPPQQVPFP HNGPSGGQGSFPGGMGFPGEGPLGRPSNLPQ SSADAALCKPGGPGGPDSFTVLGNSMPSVFT DPDLQEVIRPGATGIPEFDLSRIIPSEKPSQTLQ YFPRGEVPGRKQPQGPGPGFSHMQGMMGEQ APRMGLALPGMGGGPGPVGTPDIPLGTAPSMP
		ļ				GHNPMRPPAFLQQGMMGPHIIRMMSPAQST MPGQPTLMSNPAAAVGMIPGKDRGPAGLYT
829	2179	A	6797	433	3	HPGPVGSPGMMMSMQGMMGP\NRTS ASFFNFSICICKIILEVGPPVGHPAHDDVGGRH
023						GPGGR/GSRSPRSLQCAPGGGRRSGCPAGSSP ASTCPPSPGGSGADRFGPSPPPPSREAAPTAG AAASSTSSGASCPPVPASSRWGVRSRTRSGSG GEREPRDRPSERPRLV
830	2180	A	6800	3	1911	LPERAFGPRTPRAPRRRRRRLLLSPPPRPPPPL DREPRAPGPWLCPSRAGTAQDPARIRERRGR VAGGAAGPAMELRARGWWLLCAAAALVAC ARGDPASKSRSCGEVRQIYGAKGFSSS\DVPQ AEISGEHLRICPQGYTCCTSEMEENLANRSHA ELETALRDSSRVLQAMLATQLRSFDDHFQHL LNDSERTLQATFPGAFGELYTQNARAFRDLY SELRLYYRGANLHLEETLAEFWARLLERLFK QLHPQLLLPDDYLDCLGKQAEALRPF\GEAP\ RELRLRAT\RA\FVAAR\SFVQGLGVAS\DVVR KVAQVPLG\PEC\SRAVIEAGSYC/ALHCVGVP GARPCPDYCRNVLKGCLANQADLDAEWRNL LDSMVLITDKFWGTSGVESVIGSVHTWLAEA INALQDNRDTLTAKVIQGCGNPKVNPQGPGP EEKRRRGKLAPRERPPSGTLEKLVSEAKAQL RDVQDFWISLPGTLCSEKMALSTASDDRCWN GMARGRYLPEVMGDGLANQINNPEVEVDIT KPDMTIRQQIMQLKIMTNRLRSAYNGNDVDF QDASDDGSGSGSGDGCLDDLCGRKVSRKSSS SRTPLTHALPGLSEQEGQKTSAASCPQPPTFL LPLLLFLALTVARPRWR
831	2181	A	6808	2	1522	ASRHGMTPGALLMLLGALGPPLAPGVRGSEA EGRLREKLFSGYDSSVRPAREVGDRVRVSVG LILAQLISLNEKDEEMSTKVYLDLEWTDYRLS WDPAEHDGIDSLRITAESVWLPDVVLLNNND GNFDVALDISVVVSSDGSVRWQPPGIYRSSCS IQVTYFPFDWQNCTMVFSSYSYDSSEVSLQT GLGPDGQGHQEIHIHEGTFIENGQWENIHKPS RLIQPPGDPRGGREGQRQEVIFYLIIRRKPLFY LVNVIAPCILITLLAIFVFYLPPDAGEKMGLSIF ALLTLTVFLLLLADKVPETSLSVPIIIKYLMFT MVLVTFSVILSVVVLNLHHRSPHTHQMPLWV RQIFIHKLPLYLRLKRPKPEDLMPEPPHCSSP GSGWGRGTDEYFIRKPPSDFLFPKPNRFQPEL SAPDLRRFIDGPNRAVALLPELREVVSSISYIA RQLQEQEDHDALKEDWQFVAMVVDRLFLW TFIIFTSVGTL\VIFLDATYHLPPPDPFP
832	2182	A	6824	71	1079	ETMAKNPPENCEDCHILNAEAFKSKKICKSLK ICGLVFGILALTLIVLFWGSKHFWPEVPKKAY DMEHTFYSNGEKKKIYMEIDPVTRTEIFRSGN GTDETLEVHDFKNGYTGIYFVGLQKCFIKTQI KVIPEFSEPEEEIDENEEITTFFEQSVIWVPAE KPIENRDFLKNSKILEICDNVTMYW\INPTL\IS

<u> </u>	L COMO TO	1 5 2 .		1 50 11 1	I 70 - 41 - 1	
SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide	İ	in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence	ļ	09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence	ł		914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
ļ		i		amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
	İ	Į		peptide		/=possible nucleotide deletion, \=possible
	<u> </u>	<u> </u>		sequence		nucleotide insertion
	ļ			Ì		GTFAKQLHHNFAFIILVSELQDFEEEGEDLHFP
						ANEKKGIEQNEQWVVPQVKVEKTRHARQAS
ĺ	i	ĺ		1	ĺ	EEELPINDYTENGIEFDPMLDERGYCCIYCRR
						GNRYCRRVCEPLLGYYPYPYCYQGGRVICRV
						IMPCNWWVARMLGRV
833	2183	A	6846	116	602	EAEGEQVCGAKCCGDAPHVENREEETARIGP
i	ľ	1		ĺ	ľ	GVMESKEERALNNLIVENVNQENDEKDEKE
ļ						QVANKGEPLALPLNVSEYCVPRGNRRRFRVR
		]				QPILQYRWDIMHRLGEPQARMREENMERIGE
	i					EVRQLMEKLREKQLSHSLRAVSTDPPHHDIIII
						DEFC\LMP
834	2184	A	6851	3	2024	PNGVALLHLPGAAVIPNTNYMFQDALGGRSR
	1	ļ				GSREESPAPSRAPASASLWRRLVVVEAKMAA
						HAAAAAQAAAAQAAHAEAADSWYLALLGF
						AEHFRTSSPPKIRLCVHCLQAVFPFKPPQRIEA
	ļ					RTHLQLGSVLYHHTKNSEQARSHLEKAWLIS
						QQIPQFEDVKFEAASLLSELYCQENSVDAAKP
			1		i	LLRKAIQISQQTPYWHCRLLFQLAQLHTLEKD
						LVSACDLLGVGAEYARVVGSEYTRALFLLSK
į	1					GMLLLMERKLQEVHPLLTLCGQIVENWQGN
	1					PIQKESLRVFFLVLQVTHYLDAGQVKSVKPC
						LKOLOOCIOTISTLHDDEILPSNPADLFHWLP
			1			KEHMCVLVYLVTVMHSMQAGYLEKAQKYT
j	,	j			J	DKALMQLEKLKMLDCSPILSSFQVILLEHIIM
	}					CRLVTGHKATALQEISQVCQLCQQSPRLFSN
	i					HAAQLHTLLGLYCVSVNCMDNAEAQFTTAL
}	l	ł		ļ	}	RLTNHQELWAFIVTNLASVYIREGNRHQEVV
						LYSLLERINPDHSFPVSSHCLRAAAFYVRGLF
	1	1				SFFQGRYNEAKRFLRETLKMSNAEDLNRLTA
	ļ					CSLVLLGHIFYVLGNHRESNNMVVPAMQLAS
ļ	1	1	1			KIPDMSVQLWSSALLRDLNKACGNAMDAHE
						AAQMHQNFSQQLLQDHIEACSLPEHNLITWT
						DGPPPVQFQAQNGPNTSLASLL
835	2185	A	6855	334	1268	PTRRPILPLTSPKAISVPSPLQGKQHTLVKSCL
633	2103	A	0033	334	1200	
						SVSGIGGFLVSLSSRMKLQTLAVSVTALKFWS
						AYVPCQTQDRDALRLTLEQIDLIRRMCASYSE
1		1	1			LELVTSAKALNDTQKLACLIGVEGGHSLDNS
1		1				LSILRTFYMLGVRYLTLTHTCNTPWAESSAK
		1	1			GVHSFYNNISGLTDFGEKVVAEMNRLGMMV
		ĺ				DLSHVSDAVARRALEVSQAPVIFSHSAARGV
						CNSARNVPDDILQLLEEERWAFVMVSLFHGE
1						LIQWQPIRPMCSTVADHFDHIKAV\IGSKFIGI
						GGDYDGAGKYRKKTTCKAPWRTSSRMSS
836	2186	Α	6862	315	11	PPRSRPSCWRKKVGPGRPWWWGGTGPPGQG
1		İ				RPEIRLLPLPMTGACGAVAASRTGSSGPG/SSL
•		١.	]			PNGHGGKGSGLANGLAGNP\GHLGLGSSFGT
<u></u>			1			GPGSGRPPP
837	2187	A	6863	2	1615	VLRGQRGPAGGLAEERRRGRNEWRIHDVTT
		1				APFPGLVQRRSRLLIVSQVRYFLKNKVSPDLC
l						NEDGLTALHQCCIDNFEEIVKLLLSHGANVN
		1	-	ĺ	İ	AKDNELWTPLHAAATCGHINLVKILVOYGA
						DLLAVNSDGNMPYDLCEDEPTLDVIETCMAY
J	]		J			QGITQEKINEMRVAPEQQMIADIHCMIAAGQ
						DLDWIDAQGATLLHIAGANGYLRAAELLLDH
						GVRVDVKDWDGWEPLHAAAFWGQMQMAE
						LLVSHGAN\LNARTSMDEMPIDLCEEEEFKVL
J	)	]	]		,	LLELK\HKHDVIMKSQLRHKSSLSRRTSHRQA
					,	S/SVGKVVRRTQPVGTGPNL\YRKEYE/GEEAI
1						LWQRSA\AEDQRTSTYNGDIRET\RTDQENKD
				ı	i	L P 11 OKOK KARDOK I ST T NODJIKE I KTOVENKU

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  PNPRLEK\PVLLSEFPTKIPRGELDMPVENGLR APVSAYQYALANGDVWKVHEVPDYSMAYG NPGVADATPPWSSYKEQSPQTLLELKRQRAA AKLLSHPFLSTHLGSSMARTGESSSEGKAPLI GGRTSPYSSNGTSVYYTVTSGDPPLLKFKAPI EEMEEKVHGCCRIS
838	2188	A	6865	6291	739	AGPLEPRVQGAMALQLWALTLLGLLGAGAS LRPRKLDFFRSEKELNHLAVDEASGVVYLGA VNALYQLDAKLQLEQQVATGPVLDNKKCTP PIEASQCHEAEMTDNVNQLLLVDPPRKRLVE CGQLLKGI\CALRALSNISLRLFYEDGSGEKSF VASNDEGVATVGLVSSTOPGGDRVLFVGKG NGPHDNGIIVSTRLLDRTDSREAFEAYTDHAT YKAGYLSTNTQQFVAAFEDGPYVFFVRQQD KHPARNRTLLARMCREDPNYYSYLEMDLQC RDPDIHAAAFGTCLAASVAAPGSGRVLYAVF SRDSRSSGGPGAGLCLFPLDEVHAKMEANRN ACYTGTREARDIFYKPFHGDIQCGGHAPGSSK SFPCGSEHLPYPLGSRDGLRGTAVLQRGGLN LTAVTVAAENNHTVAFLGTSDGRILKVYLTP DGTSSEYDSILVEINKRVKRDLVLSGDLGSLY AMTQDKVFRLPVQECLSYPTCTQCRDSQDPY CGWCVVEGRCTTRKAECPRAEEASHWLWSRS KSCVAVTSAQPQNMSRRAQGEVQLTVSPLPA LSEEDELLCLFGESPPHPARVEGEAVICNSPSS IPVTPPGQDHVAVTIQLLLRRGNIFLTSYQYPF YDCRQAMSLEENLPCISCVSNRWTCQWDLR YHECREASPNPEDGIVRAHMEDSCPQFLGPSP LVIPMNHETDVNFQGKNLDTVKGSSLHVGSD LLKFMEPVTMQESGTFAFRTPKLSHDANETL PLHLYVKSYGKNIDSKLHVTLYDCSFGRSDC SLCRAANPDYRCAWCGGQSRCVYEALCNTT SECPPPVITRIQPETGPLGGGIRITILGSNLGVQ AGDIQRISVAGRNCSFQPERYSVSTRIVCVIEA AETPFTGGVEVDVFGKLGRSPPNVQFTFQQP KPLSVEPQQGPQAGGTTLTHGTHLDTGSQED VRVTLNGVPCKVTKFGAQLQCVTGPQATRG QMLLEVSYGGSPVPNPGIFTYRENPVLRAFE PLRSFASGGRSINVTQGFSLIQRFAMVVIAEP LQSWQPPREAESLQPMTVVGTDYVFHNDTK VVFLSPAVPEEPEAYNLTVLIEMDGHRALLRT EAGAFEYVPDFTFENFTGGVKKQVNKLIRAR GTNLNKAMTLQEAEAFVGAERCTMKTLTET DLYCEPPEVQPPPKRRQKRDTTHNLPEFIVKF GSREWVLGRVEYDTRVSDVPLSLILPLVIVPM VVVIAVSVYCYWRKSQQAEREYEKIKSQLEG LEESVRDRCKKEFTDLMIEMEDQTNDVHEAG IPVLDYKTYTDRVFFLPSKDGDKDVMITGKL USPEPFRPVVEQALYQFSNLLNSKSFLINFIHT L\ENQPEFSARAKVYFASLLTVALHGKLEYYT DIMHTLFLELLEQYVVAKNPKLMLRRSETVV ERMLSNWMSICLYQYLKDSAGEPLYKLFKAI KHQVEKGPVDAVQKKAKYTLNDTGLLGDD VEYAPLTVSVIVQDEGVDAIPVKVLNCDTISQ VKEKIIDQVYRGQPCSCWPRPDSVVLEWRPG STAQILSDLDLTSQREGRWKRVNTLMHYNVR DGATLILSKVGVSQQPEDSQQDLPGERHALL EEENRVWHLVRPTDEVDEGKSKRGSVKEKE RTKAITEIYLTRLLSVKGTLQQFVDNFFQSVL APGHAVPPAVKYFFDFLDEQAEKHNIQDEDTI

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion HIWKTNSLPLRFWVNILKNPHFIFDVHVHEVV DASLSVIAQTFMDACTRTEHKLSRDSPSNKLL YAKEISTYKKMVEDYYKGIRQMVQVSDQDM NTHLAEISRAHTDSLNTLVALHQLYQYTQKY YDEIINALEEDPAAQKMQLAFRLQQIAAALE NKVTDL
839	2189	A	6872	1	1485	RARRLALQCHVCVCALTPGEQSGRRLPGQT WLMFSCFCFSLQDNSFSSTTVTECDEDPVSLH EDQTDCSSLRDENNKENYPDAGALVEEHAPP SWEPQQQNVEATVLVDSVLRPSMGNFKSRKP KSIFKAESGRSHGESQETEHVVSSQSECQVRA GTPAHESPQNNAFKCQETVRL\QPRIDQRTAT SPKDAFETR\QDLNEEEAAQVHGVKDPAPAS TQSVLA\DGTDSADPSPVHKDGQNEADSAPE DLHSVGTSRLLL/YHITDGDNPTAVRHGCSL/F SGQSQRFNLDPESAPSPPSTQQFMMPRSSSRC SCGDGKEPQTITQLTKHIQSLKRKIRKFEEKFE QEKKYRPSHGDKTSNPEVLKWMNDLAKGRK QLKELKLKLSEEQGSAPKGPPRNLLCEQPTVP RENGKPEAAGPEPSSSGEETPDAALTCLKERR EQLPPQEDSKVTKQDKNLIKPLYDRYRIIKQIL STPSLIPTIVSQDTCMLLLCTDV
840	2190	A	6873	2	2054	FFRFYFSFIRLFAMSLADLTKTNIDEHFFGVAL ENNRRSAACKRSPGTGDFSRNSNASNKSVDY SRSQCSCGSLSSQYDYSEDFLCDCSEKAINRN YLKQPVVKEKEKKYNVSKISQSKGQKEISV EKKHTWNASLFNSQIHMIAQRRDAMAHRILS ARLHKIKGLKNELADMHHKLEAILTENQFLK QLQLRHLKAIGKYENSQNNLPQIMAKHQNEV KNLRQLLRKSQEKERTLSRKLRETDSQLLKT KDILQALQKLSEDKNLAEREELTHKLSIITTK MDANDKKIQSLEKQLRLNCRAFSRQLAIETR KTLAAQTATKTLQVEVKHLQQKLKEKDREL EIKNIYSHRILKNLHDTEDYPKVSSTKSVQAD RKILPFTSMRHQGTQKSDVPPL/TTKGKKATG NIDHKEKSTEINHEIPHCVNKLPKQEDSKRKY EDLSGEEKHLEVQILLENTGRQKDKKEDQEK KNIFVKEEQELPPKIIEVIHPERESNQEDVLVR EKFKRSMQRNGVDDTLGKGTAPYTKGPLRQ RRHYSFTEATENLHHGLPASGGPANAGNMR YSHSTGKHLSNREEMELEHS\DSGYEPSFGKS SRIKVKDTTFRDKKSSLMEELFGSGYVLKTD QSSPGVAKGSEEPLQSKESHPLPPSQASTSHA FGDSKVTVNSIKPSSPTEGKRKIII
841	2191	A	6874	3	2867	SSRTREMEEKEILRRQIRLLQGLIDDYKTLHG NAPAPGTPAASGWQPPTYHSGRAFSARYPRP SRRGYSSHHGPSWRKKYSLVNRPPGPSDPPA DHAVRPLHGARGGQPPVPQQHVLERQVQLS QGQNVVIKVKPPSKSGSASASGAQRGSLEEFE DTPWSDQRPREGEGEPPRGQLQPSRPTRARG TCSVEDPLLVCQKEPGKPRMVKSVGSVGDSP REPRRTVSESVIAVKASFPSALPPRTGVALG RKLGSHSVASCAPQLLGDRRVDAGHTDQPVP SGSVGGPARPASGPRQAREASLVVTCRTNKF RKNNYKWVAASSKSPRVARRALSPRVAAEN VCKASAGMANKVEKPQLIADPEPKPRKPATS SKPGSAPSKYKWKASSPSASSSSFRWQSEAG SKDHASQLSPVLSRSPSGD\RPALAHSGLKPLS GETPLSAYKVKTRTKIIRRRGSTSLPGDKKSG TSPAATAKSHLSLRRRQALRGKSSPVLKKTPN

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
						KGLVQVTKHRLCRLPPSRAHLPTKEASSLHA VRTAPTSKVIKTRYRIVKKTPASPLSAPPFPLS LPSWRARRLSLSRSLVLNRLRPVASGGGKAQ PGSPWWRSKGYRCIGGVLYKVSANKLSKTSG QPSDAGSRPLLRTGRLDPAGSCSRSLASRAVQ RSLAIIRQARQRREKRKEYCMYYNRFGRCNR GERCPYHDPEKVAVCTRFVRGTCKKTDGTC PFSHHVSKEKMPVCSYFLKGICSNSNCPYSHV YVSRKAEVCSDFLKGYCPLGAKCKKHTLLC PDFARRGACPRGAQCQLLHRTQKRHSRRAAT SPAPGPSDATARSRVSASHGPRKPSASQRPTR QTPSSAALTAAAVAAPPHCPGGSASPSSSKAS SSSSSSSSPPASLDHE\APSLQEAALAAACSNR LCKLPSFISLQSSPSPGAQPRVRAPRAPLTKDS GKPLHIKPRL
842	2192	A .	6898	506	2071	WPDLVHTWSSEEAMGSCCSCPDKDTVPDNH RNKFKVINVDDDGNELGSGIMELTDTELILYT RKRDSVKWHYLCLRRYGYDSNLFSFESGRRC QTGQGIFAFKCARAEELFNMLQEIMQNNSIN VVEEPVVERNNHQTELEVPRTPRTPTTPGFAA QNLPNGYPRYPSFGDASSHPSSRHPSVGSARL PSVGEESTHPLLVAEEQVHTYVNTTGVQEER KNRTSVHVPLEARVSNAESSTPKEEPSSIEDR DPQILLEPEGVKFVLGPTPVQKQLMEKEKLE QLGRDQVSGSGANNTEWDTGYDSDERRDAP SVNKLVYENINGLSIPSASGVRRGRLTSTSTSD TQNINNSAQRTALLNYENLPSLPPVWEARK LSRDEDDNLGPKTPSLNGYHNNLDPMHNYV NTENVTVPASAHKIEYSRRRDCTPTVFNFDIR RPSLEHRQLNYIQVDLEGGSDSDNPQTPKTPT TPLPQTPTRRTELYAVIDIERTAAMSNLQKAL PRDDGTSRKTRHNSTVDLPL
843	2193	A	6919	2	663	AGRPGTTHASGKMAYQSLRLEYLQIPPVSRA YTTACVLTTAAVQLELITPFQLYFNPELIFKHF QIWRLITNFLFFGPVGFNFLFNMIFLYRYCRM LEEGSFRGRTADFVFMFLFGGFLMTLFGLFVS L/VFLGPGLYNN/GSSMCGAE\EPLCPHELLRP SQLPGPLSALGAHGIFLVVGELNHCGPFGYCS WTHIFFLGRCISQSTWWNKNSENTIYFESYF
844	2194	A	6928	902	366	HRLCMPIQGACGERME/FSLLLPGLECNGVIL AHCNLRLPGSSNSPASASQVAGITGVCHHAR LIFVFSVETGFLHAGQAGLELLTSGDPPASAS QSAGITGKSQHTRPGYEFIIPYSAAQEDALKA LM
845	2195	A	6939	1660	317	LYPENLGESLFPILLLPPPWPDGGRPCCVEMS TRAKKLRRIWRILEEKESVAGAVQTLLLRSQE GGV\TSAAASTLSEPPRRTQESRTRTRALGLPT LPMEKLAASTEPQGPRPVLGRESVQVPDDQD FRSFRSECEAEVGWNLTYSRAGVSVWVQAV EMDRTLHKIKCRMECCDVPAETLYDVLHDIE YRKWDSNVIETFDIARLTVNADVGYYSWR CPKPLKNRDVITLRSWLPMGADYIIMNYSVK HPKYPPRKDLVRAVSIQTGYLIQSTGPKSCVIT YLAQVDPKGSLPKWVVNKSSQFLAPKAMKK MYKACLKYPEWKQKHL\PHFKPWL\HPEQSP LPSLALS\ELSVQHADS\LENIDESAV\AESREE R\MGGAGGEG\SDDDTSLYAEAPHRFRETETG PGAGRALGAAAAPALSPLHPPGTWWHRARP RRVLQPGWTEPQ

r on one	1 000	132	Tana	1 72		
SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence	İ		914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
}	ļ		ł	amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
	ļ			residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
]	j	J	]	peptide	]	/=possible nucleotide deletion, \=possible
		<u> </u>	<del> </del>	sequence		nucleotide insertion
846	2196	A	6944	42	2672	RRKMAGCRGSLCCCCRWCCCCGERETRTPE
		ĺ			ĺ	ELTILGETQEEEDEILPRKDYESLDYDRCINDP
						YLEVLETMDNKKGRRYEAVKWMVVFAIGV
1	ł	ľ	ł	ľ		CTGLVGLFVDFFVRLFTQLKFGVVQTSVEECS
		1				QKGCLALSLLELLGFNLTFVFLESLLGLIEPVE
1		l			ł	AGSGITEGKCYLYARQVPGLVRLPTLLWKAL
						GVLLTVAAMLLI\GLGSPMIHSGSVVGAGLPQ
J	J	ļ	J		1	FQSISLRKIQFNFPYFRSDRYGK\DKRDFVSAG
		ŀ	1			AAAGVAAAFGAPIGGTLFSLEEGSSFWNOGL
1						TWKVLFCSMSATFTLNFFRSGIQFGSWGSFQL
i		1			1	PGLLNFGEFKCSDSDKKCHLWTAMDLGFFV
						VMGVIGGLLGATFNCLNKRLAKYRMRNVHP
1	1					KPKLVRVLESLLVSLVTTVVVFVASMVLGEC
						RQMSSSSQIGNDSFQLQVTEDVNSSIKTFFCP
1	1	1				NDTYNDMATLFFNPQESAILQLFHQDGTFSPV
						TLALFFVLYFLLACWTYGISVPSGLFVPSLLC
	1	]			J	GAAFGRLVANVLKSYIGLGHIYSGTFALIGAA
						AFLGGVVRMTISLTVILIEST\NEITYGLPIMVT
						LMVGKWTGDFFNKGI\YDIHVGLRGVPLLEW
	1					ETEVEMDKLRASDIMEPNLTYVYPHTRIQSLV
						SILRTTVHHAFPVVTENRGNEKEFMKGNQLIS
ł	ł	l	ł			
	1					NNIKFKKSSILTRAGEQRKRSQSMKSYPSSEL RNMCDEHIASEEPAEKEDLLQQMLERRYTPY
}		ļ				
						PNLYPDQSPSEDWTMEERFRPLTFHGLILRSQ
	İ					LVTLLVRGVCYSESQSSASQPRLSYAEMAED
	1	ĺ				YPRYPDIHDLDLTLLNPRMIVDVTPYMNPSPF
'						TVSPNTHVSQVFNLFRTMGLRHLPVVNAVGE
847	2197	A	6951	3	1994	IVGIITRHNLTYEFLQARLRQHYQTI
047	2197	^	0931	3	1994	NTNSSSVTNSAAGVEDLNIVQVTVPDNEKER
ł	ł	}	1			LSSIEKIKQLREQVNDLFSRKFGEAIGVDFPVK
						VPYRKITFNPGCVVIDGMPPGVVFKAPGYLEI
ļ	ļ	ļ	Į l			SSMRRILEAAEFIKFTVIRPLPGLELSNGEYST
	İ					VGKRKIDQEGRVFQEKWERAYFFVEVQNIST
						CLICKRSMSVSKEYNLRRHYQTNHSKHYDQY
		ĺ				MERMRDEKLHELKKGLRKYLLGLSDTECPE
						QKQVFANPSPTQKSPVQPVEDLAGNLWEKLR
1			[		l	EKIRSFVAYSIAIDEITDINNTTQLAIFIRGVDE
		}				NFDVSEELLDTVPMTGTKSGNEIFSRVEKSLK
1	1			·	l	NFCINWSKLVSVASTGTPPMVDANNGLVTKL
				ļ		KSRVATFCKGAELKSICCIIHPESLCAQ\KLKM
			] ]	}	J	DHVMDVVVKSVNWICSRGLNHSEFTTLLYEL
						DSQYGSLLYYTEIKWLSRGLVLKRFFESLEEI
				l		DSFMSSRGKPLPQLSSIDWIRDLAFLVDMTM
				l		HLNALNISLQGHSQIVTQMYDLIRAFLAKLCL
	}		1 1	ł		WETHLTRNNLAHFPTLKLVSRNESDGLNYIP
					l	KIAELKTEFQKRLSDFKLYESELTLFSSPFSTKI
1			, ,	j	ļ	DSVHEELQMEVIDLQCNTVLKTKYDKVGIPE
						FYKYLWGSYPKYKHHCAKILSMFGSTYICEQ
<u></u>		L				LFSIMKLSKTKYCSQLKDSQWDSVLHIAT
848	2198	A	6985	3	289	SVQYLPGRPTRTHASTDAPLMLKFTPLPSKTK
						ASAPVQCLLLMAATFSPQGLAKPHSGTIPIT\C
			( l	İ		CFNAINTKIPIQRLESYTRITNIQCPKEAVM
849	2199	A	6999	963	5	LDFLCHRDMGDNITSITEFLLLGFPVGPRIQM
]	1 4199				-	LLFGLFSLFYVFTLLGNGTILGLISLDSRLHAP
	2199		1 1			THE CHI CHI I VI INCOMMENCE
	2199				I	MVEEL SHI / AVVDIAV A CNITTIOD NOT VIXIT T TITE
	2199					MYFFLSHL\AVVDIAYACNTVPRMLVNLLHP
	2199					AKPISFAGRMMQTFLFSTFAVTECLLLVVMS
	2199					AKPISFAGRMMQTFLFSTFAVTECLLLVVMS YDLYV\AICHPLRYLAIMTWRVCITLAVTSWT
	2199					AKPISFAGRMMQTFLFSTFAVTECLLLVVMS YDLYV\AICHPLRYLAIMTWRVCITLAVTSWT TGVLLSLIHLVLLLPLPFCRPQKIYHFFCEILA
	2199					AKPISFAGRMMQTFLFSTFAVTECLLLVVMS YDLYV\AICHPLRYLAIMTWRVCITLAVTSWT

	,					
SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline.
uence		l	914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
40	ļ		**.	amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
			1		sequence	
				peptide		/=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion
						GLFYGTAIIMYVGPRYGNPKEQKKYLLLFHS
	1	ļ				LFNPMLNPLICSLRNSEVKNTLKRVLGVERAL
850	2200	A	7001	1	1011	MGNDSVSYEYGDYSDLSDRPVDCLDGACLAI
			1	_		DPLRVAPLPLYAAIFLVGVPGNAMVAWVAG
						KVARRRVGATWLLHLAVADLLCCLSLPILAV
		Į				PIARGGHWPYGAVGCRALPSIILLTMYASVLL
		1				LAALSADLCFLALGPAW\CLRFS/GACGVQVA
ļ	ł	į	1		ļ	CGAAWTLALLLTVPSAIYRRLHQEHFPARLQ
						CVVDYGGSSSTENAVTAIRFLFGFLGPLVAVA
						SCHSALLCWAARRCRPLGTAIVVGFFVCWAP
		İ			·	YHLLGLVLTVAAPNSALLARALRAEPLIVGL
			1			ALAHSCLNPMLFLYFGRAQLRRSLPAACHW
	J	J	1	]		ALRESOGODESVDSKKSTSHDLVSEMEV
051	2201	1	7011	1	2210	
851	2201	A	7011	1	2310	AAASPLRMSRKGPRAEVCADCSAPDPGWASI
		]				SRGVLVCDECCSVHRSLGRHISIVKHLRHSA
	1	l	1		}	WPPTLLQMVHTLASNGANSIWEHSLLDPAQV
			i			QSGPALKQTPKDKV\HPIKSEFIRAKYQMLAF
						VHKLPCRDDDGVTAKDLSKQLHSSVRTGNLE
		ļ				TCLRLLSLGAQANFFHPEKGTTPLHVAAKAG
		1				QTLQAELLVVYGADPGSPDVNGRTPIDYARQ
Į.	1	1	1		į	AGHHELAERLVECQYELTDRLAFYLCGRKPD
		İ				~
1			i			HKNGHYIIPQMADSLDLSELAKAAKKKLQAL
						SNRLFEELAMDVYDEVDRRENDAVWLATQN
						HSTLVTERSAVPFLPVNPEYSATRNQGRQKL
ļ	1		]			ARFNAREFATLIIDILSEAKRRQQGKSLSSPTD
						NLELSLRSQSDLDDQHDYDSVASDEDTDQEP
			1			LRSTGATRSNRARSMDSSDLSDGAVTLQEYL
	1		İ			ELKKALATSEAKVQQLMKVNSSLSDELRRLQ
						REIHKLQAENLQLRQPPGPVPTPPLPSERAEH
}						TPMAPGGSTHRRDRQAFSMYEPGSALKPFGG
			1			PPGDELTTRLQPFHSTELEDDAIYSVHVPAGL
						YRIRKGVSASAVPFTPSSPLLSCSQEGSRHTSK
1		l	ł			LSRHGSGADSDYENTQSGDPLLGLEGKRFLE
						LGKEEDFHPELESLDGDLDPGLPSTEDVILKT
						EQVTKNIQELLRAAQEFKHDSFVPCSEKIHLA
						VTEMASLFPKRPALEPVRSSLRLLNASAYRLQ
			ł			SECRKTVPPEPGAPVDFQLLTQQVIQCAYDIA
1		ł	1	}	1	KAAKQLVTITTREKKQ
8.52	2202	A	7016	484	1777	RISKIQVYYSTGYSSRKMNPTLGLAIFLAVLL
2.0.0	2202	۱ ^	/ / / /	704	1///	
ĺ		1				TVKGLLKPSFSPRNYKALSEVQGWKQRMAA
						KELARQNMDLGFKLLKKLAFYNPGRNIFLSP
1	1	l	}	}		LSISTAFSMLCLGAQDSTLDEIKQGFNFRKMP
	1					EKDLHEGFHYIIHELTQKTQDLKLSIGNTLFID
						QRLQPQRKFLEDAKNFYSAETILTNFQNLEM
		ļ				AQKQINDFI/ESKTHGKINNLIENIDPGTVMLL
1	ł	ł	}			ANYIFFRARWKHEFDPNVTKEEDFFLEKNSS
	1	ĺ	l '			VKVPMMFRSGIYQVGYDDKLSCTILEIPYQK
1			1			NITAIFILPDEGKLKHLEKGLQVDTFSRWKTL
1						
		1				LSRRVVDVSVPRLHMTGTFDLKKTLSYIGVS
1	ł	{	}	1		KIFEEHGDLTKIAPHRSLKVGEAVNKAELKM
1	1	1				DERGTEGAAGTGAQTLPMETPLVVKIDKPYL
						LLIYSEKIPSVLFLGKIVNPIGK
853	2203	A	7017	1	3293	MTHACNPSTLGGQGRRITRSHGRRRSSRGPV
						ARHVAAGAGHENKHGGSRRFPAGVAPRRAM
ļ	1	ļ	) .			ANVSKKVSWSGRDRDDEEAAPLLRRTARPG
		ļ				
						GGTPLLNGAGPGAARQSPRSALFRVGHMSSV
	1	l				ELDDELLEP\DMDPPHPFPKEIPHNEKLLSLKY
	1	1				ESLDYDNSENQLFLEEERRINHTAFRTVEIKR
1	ļ	J	J			WVICALIGILTGLVACFIDIVVENLAGLKYRVI
		<u></u>	1			KGSILPNIDKFTEKGGLSFSLLLWATLNAAFV

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  LVGSVIVAFIEPVAAGSGIPQIKCFLNGVKIPH  VVRLKTLVIKVSGVILSVVGGLAVGKEGPMI  HSGSVIAAGISQGRSTSLKRDFKIFEYFRRDTE  KRDFVSAGAAAGVSAAFGAPVGGVLFSLEEG  ASFWNQFLTWRIFFASMISTFTLNFVLSIYHG  NMWDLSSPGLINFGRFDSEKMAYTIHEIPVFI  AMGVVGGVLGAVFNALNYWLTMFRIRYHR  PCLQVIEAVLVAAVTATVAFVLIYSSRDCQPL  QGGSMSYPLQLFCADGEYNSMAAAFFNTPEK  SVVSLFHDPPGSYNPLTLGLFTLVYFFLACWT  YGLTVSAGVFIPSLLIGAAWGRLFGISLSYLTG  AAIWADPGKYALMGAAAQLGGIVRMTLSLT  VIMMEATSNVTYGFPIMLVLMTAKIVGDVFIE  GLYDMHIQLQSVPFLHWEAPVTSHSLTAREV  MSTPVTCLRRREKVGVIVDVLSDTASNHNGF  PVVEHADDTQPARLQGLILRSQLIVLLKHKVF  VERSNLGLVQRRLRLKDFRDAYPRFPPIQSIH  VSQDERECTMDLSEFMNPSPYTVPQEASLPR  VFKLFRALGLRHLVVVDNRNQVVGLVTRKD  LARYRLGKRGLEELSLAQTGPKAQATAEGRV  AGAAQQPCQLRAVTLEDLGLLLAGGLASPEP  LSLEELSERYESSHPTSTASVPEQDTAKHWNQ  LEQWVYELQAEVACLREHKQRCERATRSLL  RELLQVRARVQLQGSELRQLQQEARPAAQAP  EKEAPEFSGLQNQMQALDKRLVEVREALTRL
854	2204	A	7037	139	2604	QEEQGREVACGALQKNQEDSSRRVDLEVAR M  AGTWEPRPYDQAKETGAPGSQPPVPPMELRP WLLWVVAATGTLVLLAADAQGQKVFTNTW AVRIPGGPAVANSVARKHGFLNLGQIFGDYY HFWHRGVTKRSLSPHRPRHSRLQREPQVQWL EQQVAKRRTKRDVYQEPTDPKFPQQWYL\SG VTQ\RDLMVKAAWAQGYTGHGIVVSILDDGI EKNHPDLAGNYDPGASFDVNDQDPDPQPRY TQMNDNRHGTRCAGEVAAVANNGVCGVGV AYNARIGGVRMLDGEVTDAVEARSLGLNPN HIHIYSASWGPEDDGKTVDGPARLAEEAFFR GVSQGRGGLGSIFVWASGNGGREHDSCNCD GYTNSIYTLSISSATQFGNVPWYSEACSSTLA TTYSSGNQNEKQIVTTDLRQKCTESHTGTSAS APLAAGIIALTLEANKNLTWRDMQHLVVQTS KPAHLNANDWATNGVGRKVSHSYGYGLLD AGAMVALAQNWTTVAPQRKCIIDILTEPKDI GKRLEVRKTVTACLGEPNHITRLEHAQARLT LSYNRRGDLAIHLVSPMGTRSTLLAARPHDY SADGFNDWAFMTTHSWDEDPSGEWVLEIEN TSEANNYGTLTKFTLVLYGTAPEGLPVPPESS GCKTLTSSQACVVCEEGFSLHQKSCVQHCPP GFAPQVLDTHYSTENDVETIRASVCAPCHAS CATCQGPALTDCLSCPSHASLDPVEQTCSRQS QSSRESPPQQPPRLPPEVEAGQRLRAGLLPS HLPEVVAGLSCAFIVLVYTVFLVLQLRSGFS FRGVKVYTMDRGLISYKGLPPEAWQEECPSD SEEDEGRGERTAFIKDQSAL ORPASOLLAPFAAEALPGAPRAAMAOHFSLA
855	2205	A	7058	3	1441	QRPASQLLAPFAAEALPGAPRAAMAQHFSLA ACDVVGFDLDHTLCRYNLPESAPLIYNSFAQF LVKEKGYDKELLNVTPEDWDFCCKGLALDL EDGNFLKLANNGTVLRASHGTKMMTPEVLA EAYGKKEWKHFLSDTGMACRSGKYYFYDN

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \text{\text{=}possible} nucleotide insertion  YFDLPGALLCARVVDYLTKLNNGQKTFDFW KDIVAAIQHNYKMSAFKENCGIYFPEIKRDPG RYLHSRPESVKKWLRQLKNAGKILLLITSSHS DYCRLLCA\YILGNDFTDLFDIVITNALKPGFF SHLPSQRPFRTLENDEEQEALPSLDKPGWYSQ GNAVHLYELLKKMTGKPEPKVVYFGDSMHS DIFPARHYSNWETVLILEELRGDEGTRSQRPE ESEPLEKKGKYEGPKAKPLNTSSKKWGSFF\I DSVLGLENTEDSLVYTWSCKRISTYSTIAIPSI EAIAELPLDYKFTRFSSSNSKTAGYYPNPPLV
856	2206	A	7082	396	1635	LSSDETLISK  SSPSVFEFEHAVQPVFTMEFLKTCVLRRNACT AVCFWRSKVVQKPSVRRISTTSPRSTVMPAW VIDKYGKNEVLRFTQNMMMPIIHYPNEVIVK VHAASVNPIDVNMRSGYGATALNMKRDPLH VKIKGEEFPLTLGRDVSGVVMECGLDVKYFK PGDEVWAAVPPWKQGTLSEFVVVSGNEVSH KPKSLTHTQAASLPYVALTAWSAINKVGGLN DKNCTGKRVLILGASGGVGTFAIQVMKAWD AHVTAVCSQDASELVRKLGADDVIDYKSGSV EEQLKSLKPFDFILDNVGGSTETWAPDFLKK WSGATYVTLVTPFLLNMDRLGIADGMLQTG VTVGSKALKHFWKGVHYRWAFFMASGPCL DDIAELVDAGKIRPVIEQTFPFSKVPEAFLKV ERGHARGKTVINVV
857	2207	A	7088	320	2417	LRRRKMTPQSLLQTTLFLLSLLFLVQGAHGR GHREDFRFCSQRNQTHRSSLHYKPTPDLRISIE NSEEALTVHAPFPAAHPASRSFPDPRGLYHFC LYWNRHAGRLHLLYGKRDFLLSDKASSLLCF QHQEESLAQGPPLLATSVTSWWSPQNISLPSA ASFTFSFHSPPHTGAHNASVDMCELKRDLQL LSQFLKHPQKASRRPSAAPASQQLQSLESKLT SVRFMGDMGSFEEDRINATVWKLQPTAGLQ DLHIHSRQEEEQSEIMEYSVLLPRTLFQRTKG RSGEAEKRLLLVDFSSQALFQDKNSSQVLGE KVLGIVVQNTKVANLTEPVVLTFQHQLQPKN VTLQCVFWVEDPTLSSPGHWSSAGCETVRRE TQTSCFCNHLTYFAVLMVSSVEVDAVHKHY LSLLSYVGCVVSALACLVTIAAYLCSRVPLPC RRKPRDYTIKVHMNLLLAVFLLDTSFLLSEPV ALTGSEAGCRASAIFLHFSLLTCLSWMGLEG YNLYRLVVEVFGTYVPGYLLKLSAMGWGFPI FLVTLVALVDVDNYGPIILAVHRTPEGVIYPS MCWIRDSLVSYITNLGLFSLVFLFNMAMLAT MVVQILRLRPHTQKWSHVLTLLCLSLVLG\LP WALIFFSFASGTFQLVVLYLFSIITSFQGFLIFI WYWSMRLQARGGPSPLKSNSDSARLPISSGS TSSSRI
858	2208	A	7091	185	415	DAGAVKSSDTNIWFRGMCDDKKGHRCPS*G QPQHFHVAFHTEAEGAMFYFRLHVIHRVMQS QQQLFPSTLFSWLLE
859	2209	A	7136	3	302	FFFWRQSLALLPRLECSGATGAHCNLHFPGSS DCPTSAS*IAGITGACYHAWLLFVFLAETGFH HVGQGGLELLTSSDPSGSASQSAGITGVSHCT WPI
860	2210	A	7156	23	591	ALSTETRTPDMRRLLLVTSLVVVLLWEAGAV PAPKVPIKMQVKHWPSEQDPEKAWGARVVE PPEKDDQLVVLFPVQKPKLLTTEEKPRGQGR GPILPGTKAWMETEDTLGRVLSPEPDHDSLY

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0.51	0011			1200		DHIYHPQ*GSRGHHCPRPVPRPRLLGLGPSLP CPS
861	2211	A	7161	1220	1003	NYVCTIAF*EKKMGF*LSLSCLVLLFVLFLDCI LTTTTRIMFHCTYLFASVCLSLLNTLLSPNCL KSAMILQ .
862	2212	A	7211	665	847	LKYYHITMGIYKTGKKVIL*KSSMSNRFSVIF YKNIQKLSFSNYVYHQNYVFSSDWSYDF
863	2213	A	7212	924	1273	HGSSCALGDLAPG*LPSGPVLSSPAVRL*RKP LVWDSPSCLPATGPT*GLVLVLGGPDCT*WA RGQHEHKRMRAP*SCRVTVNLAKKKKKTDQ CIKPNYQSPPKECDYNILANSVA
864	2214	A	7214	845	1619	SDKGGKKADRKNHLRHAFPLLPHRVRERLH DPKVPVDADHVQGQDPGRAAHDIHGEDVTE KVSKDPLAPDEVGDTDEGHDRHGHREVGQR HGHDQEEVAYEERACEGGKFATVEVTDKPV DEALREAMPKVAKYAGGTNDKGIGMGMTV PISFAVFPNEDGSLQKKLKVWFRIPNQFQSDP PAPSDKSVKIEEREGITVYSMQFGGYAKEAD YVAQATRLRAALEGTATYRGDIYFCTGYDPP MKPYGRRNEIWLLKT
865	2215	A	7246	559	682	RRLGAVAHAYTSSTLGGRGGWIT*GQELQTS LANMAKPRLY
866	2216	A	7257	641	1310	TCTYKYLMGWIRGRRSRHSWEMSEFHNYNL DLKKSDFSTRWQKQRCPVVKSKCRENASPFF FCCFIAVAMGIRFIIMVAIWSAVFLNSLFNQEV QIPLTESYCGPCPKNWICYKNNCYQFFDESKN WYESQASCMSQNASLLKVYSKEDQDLLKLV KSYHWMGLVHIPTNGSWQWEDGSILSPNLLT IIEMQKGDCALYASSFKGYIENCSTPNTYICM ORTV
867	2217	A	7288	151	396	SIKIIEAFGSNGPDFWFFRYWSP*LFRQQVVFI MPFFQTLWLMNANRFCSIFTTTNVANNCWW TPYHCWLSVVVCRCESHGI
868	2218	A	7298	3	272	PDTVIGGRGSGGKEFGRWVLW*VFE*RLGTP KGSCPAGGSRMVSESD*EGRGC*ASYPCAC* AGS*WR*GSRPAGRGTPPRSLSHARPP
	2219	A	7332	1223	332 ,	PRRDAEDRDESCLNPAFPIGLLHPNSVNSMAR FLTLCTWLLLLGPGLLATVRAECSQDCATCS YRLVRPADINFLACVMECEGKLPSLKIWETC KELLQLSKPELPQDGTSTLRENSKPEESHLLA KRYGGFMKRYGGFMKKMDELYPMEPEEEA NGSEILAKRYGGFMKKDAEEDDSLANSSDLL KELLETGDNRERSHHQDGSDNEEEVSKRYGG FMRGLKRSPQLKEKAKELQKRYGGFMRRVG PQKW*MTSPQNRYGGFLKRFAEALPSDEEGE SYSKEVPEMEKRYGGFMRF
870	2220	A	7382	216	1018	EIHQRLTERTQFLDESRKNPNS*QANLLRGGG AGQGRGREGAESGGSRGEGPGSDGRLPATGD FWSPRSQRRGCCGRRAPRPEAMENGAVYSPT TEEDPGPARGPRSGLAAYFFMGRLPLLRRVL KGLQLLLSLLAFICEEVVSQCTLCGGLYFFEF VSCSAFLLSLLILIVYCTPFYERVDTTKVKSSD FYITLGTGCVFLLASIIFVSTHDRTSAEIAAIVF GFIASFMFLLDFITMLYEKRQESQLRKPENTT RAEALTEPLNA
871	2221	Α	7403	3	393	SCAMCSGLL*LLLPIWLSWTLGTRGSEPRSVN DPGNMSFVKETVDKLLTGFRCFREREAAPRR ALRGAALPGESEAGDPESLRSSVNADWIQYS

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872	2222	A	7413	1061	359	PFIC  FVDIVSVVEFPHCPEARFPAQHGQDSKRLTLC  PGGS*PQATLHLDRMRVSASPTKEIQVKKYK  CGLIKPCPANYFAFKICSGAANVVGPTMCFED  RMIMSPVKNNVGRGLNIALVNGTTGAVLGQ  KAFDMYSGDVMHLVKFLKEIPGGALVLVAS  YDDPGTKMNDESRKLFSDLGSSYAKQLGFRD  SWVFIGAKDLRGKSPFEQFLKEQPQTQNKYE  GWPELLEMEGCMPPKPF
873	2223	A	7429	2242	2394	ILKCAGHGGSCL*SQHFGRLRWEDRLRLGVQ DHPGOHCETPSLLKIERKLF
874	2224	A	7468	146	894	PCTSCVLWATLHLPASTRKAPQAECGMISITE WQKIGVGITGFGIFFILFGTLLYFDSVLLAFGN LLFLTGLSLIIGLRKTFWFFFQRHKLKGTSFLL GGVVIVLLRWPLLGMFLETYGFFSLFKGFFPV AFGFLGNVCNIPFLGALFRRLQGTSSMV*KTE MSSLNLDHWLKGAKREEWEPPPQSPALTHSP TYPGPPQVQKERNGAEQLTSNPQVDSRGCQE AEMQTPRRLGWGWYHTLTLYLWEEK
875	2225	A	7498	91	251	GEKPVPTWLQDEAGQWLLGFVAQPWGWPG SERHEP*HGGVLFRLGPSAPPGKL
876	2226	A	7544	403	587	YSCLCFLFKHITSFKNSVHIWLGTVVHAYNPN ILGGQGGWIA*GQEFKTSLGNTVRPCLYK
877	2227	A	7566	2	940	GCAPDTRFFVPEPGGRGAAPWVALVARGGC TFKDKVLVAARRNASAVVLYNEERYGNITLP MSHAGTGNIVVIMISYPKGREILELVQKGIPV TMTIGVGTRHVQEFISGQSVVFVAIAFITMMII SLAWLIFYYIQRFLYTGSQIGSQSHRKETKKVI GQLLLHTVKHGEKGIDVDAENCAVCIENFKV KDIIRILPCKHIFHRICIDPWLLDHRTCPMCKL DVIKALGYWGEPGDVQEMPAPESPPGRDPAA NLSLALPDDDGSDESSPPSASPAESEPQCDPSF KGDAGENTALLEAGRSDSRHGGPIS
878	2228	A	7586	315	1232	ERSLLCKVDVRWIYVSEGTKTQRRHRQGSLR RGRMQAACWYVLFLLQPTVYLVTCANLTNG GKSELLKSGSSKSTLKHIWTESSKDLSISRLLS QTFRGKENDTDLDLRYDTPEPYSEQDLWDW LRNSTDLQEPRPRAKRRPIVKTGKFKKMFGW GDFHSNIKTVKLNLLITGKIVDHGNGTFSVYF RHNSTGQGNVSVSLVPPTKIVEFDLAQQTVID AKDSKSFNCRIEYEKVDKATKNTLCNYDPSK TCYQEQTQSHVSWLCSKPFKVICIYISFYSTD YKLVQKVCPDYNYHSDTPYFPSG
879	2229	A	7605	479	391	TESWKLKWWSPTCLDQLNGSAPGNVFIHG
880		A	7612	93	659	DAAVAMTAQGGLVANRGRRFKWAIELSGPG GGSRGRSDRGSGQGDSLYPVGYLDKQVPDTS VQETDRILVEKRCWDIALGPLKQIPMNLFIMY MAGNTISIFPTMMVCMMAWRPIQALMAISAT FKMLESSSQKFLQGLVYLIGNLMGLALAVYK CQSMGLLPTHASDWLAFIEPPERMEFSGGGL LL
881	2231	A	7615	291	1452	SPQKTMRSHTITMTTTSVSSWPYSSHRMRFIT NHSDQPPQNFSATPNVTTCPMDEKLLSTVLTT SYSVIFIVGLVGNIIALYVFLGIHRKRNSIQIYL LNVAIADLLLIFCLPFRIMYHINQNKWTLGVIL CKVVGTLFYMNMYISIILLGFISLDRYIKINRSI QQRKAITTKQSIYVCCIVWMLALGGFLTMIIL TLKKGGHNSTMCFHYRDKHNAKGEAIFNFIL

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882	2232		7617		270	QLNVSSCYWKEIVHKTNEIMLVLSSFNSCLDP VMYFLMSSNIRKIMCQLLFRRFQGEPSRSEST SEFKPGYSLHDTSVAVKIQSSSKST
		A	7617	67	379	RQMALLKANKDLISAGLKEFSVLLNQQVFND PLVSEEDMVTVVEDWMNFYINYYRQQVTGE PQERDKALQELRQELNTLANPFLAKYRDFLK SHELPSHPPPSS
883	2233	A	7622	400	215	KVKTCRYNPKYSAANDTGFVDIPSREKDLAK
884	2234	A	7638	2640	2861	AVATVGPISVAVGASHVFFQFYKKGKHLSS  APVLILQMVKLSIVLTPQFLSHDQGQLTKELQ QHVKSVTCPCEYLRKVSECRQMGPGALEQFP GLSCHTSHSG
885	2235	A	7642	201	455	PSRGKMELEAMSRYTSPVNPAVFPHLTVVLL AIGMFFTAWFFVYEVTSTKYTRDIYKELLISL VASLFMGFGVLFLLLWVGIYV
886	2236	A	7692	61	569	APENPFSRQHFNSETKVKLSLKTGTWLGNHA HLGEHFSTHHELGLSGKVVGFLVKNILEVIRN GGMETRHPGKVSSWFHRWDSRAEQHNHAE HHEDVPQGDEDSKVSEAQQEFPDVVTCAGLP GLLPKALRVLLFQLKVQHRPGIHQQRPEQQD VSDHRYGRSVRQNRK
887	2237	A	7693	85	315	NPGCCLPVAMRTSYLLLFTLCLLLSEMASGG NFLTGLGHRSDHYNCVSSGGQCLYSACPIFTK IQGTCYRGKAKCCK
888	2238	A .	7702	242	1298	APSHRRRYLSPSRSAGQLGNMALERLCSVLK VLLITVLVVEGIAVAQKTQDGQNIGIKHIPAT QCGIWVRTSNGGHFASPNYPDSYPPNKECIYI LEAAPRQRIELTFDEHYYIEPSFECRFDHLEVR DGPFGFSPLIDRYCGVKSPPLIRSTGRFMWIKF SSDEELEGLGFRAKYSFIPDPDFTYLGGILNPIP DCQFELSGADGIVRSSQVEQEEKTKPGQAVD CIWTIKATPKAKIYLRFLDYQMEHSNECKRNF VAVYDGSSSIENLKAKFCSTVANDVMLKTGI GVIRMWADEGSRLNRFRMLFTSFGGASPAQA ALSFCHSNMCINNSLVCNGVQNCAYPWDEN HC
889	2239	A	7707	185	2911	CHYIMNPSTHHPASAGGSILGLFDFFGLGLGE MTMDALLARLKLLNPDDLREEIVKAGLKCGP ITSTTRFIFEKKLAQALLEQGGRLSSFYHHEA GVTALSQDPQRILKPAEGNPTDQAGFSEDRDF GYSVGLNPPEEEAVTSKTCSVPPSDTDTYRAG ATASKEPPLYYGVCPVYEDVPARNERIYVYE NKKEALQAVKMIKGSRFKAFSTREDAEKFAR GICDYFPSPSKTSLPLSPVKTAPLFSNDRLKDG LCLSESETVNKERANSYKNPRTQDLTAKLRK AVEKGEEDTFSDLIWSNPRYLIGSGDNPTIVQ EGCRYNVMHVAAKENQASICQLTLDVLENP DFMRLMYPDDDEAMI.QKRIRYVVDLYLNTP DKMGYDTPLHFACKFGNADVVNVLSSHHLI VKNSRNKYDKTPEDVICERSKNKSVELKERIR EYLKGHYYVPLLRAEETSSPVIGELWSPDQTA EASHVSRYGGSPRDPVLTLRAFAGPLSPAKAE DFRKLWKTPPREKAGFLHHVKKSDPERGFER VGRELAHELGYPWVEYWEFLGCFVDLSSQE GLQRLEEYLTQQEIGKKAQQETGEREASCRD KATTSGSNSISVRAFLDEDDMSLEEIKNRQNA ARNNSPPTVGAFGHTRCSAFPLEQEADLIEAA

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890	2240	A	7711	360	269	RHMPVIPALWEAEVGGLLEPRSSRSAWATE
891	2241	A	7721	61	1175	KLPWEPSFLIKMQIIRHSEQTLKTALISKNPVL VSQYEKLDAGEQRLMNEAFQPASDLFGPITL HSPSDWITSHPEAPQDFEQFFSDPYRKTPSPN KRSIYIQSIGSLGNTRIISEEYIKWLTGYCKAYF YGLRVKLLEPVPVSVTRCSFRVNENTHNLQIH AGDILKFLKKKKPEDAFCVVGITMIDLYPRDS WNFVFGQASLTDGVGIFSFARYGSDFYSMHY KGKVKKLKKTSSSDYSIFDNYYIPEITSVLLLR SCKTLTHEIGHIFGLRHCQWLACLMQGSNHL EEADRRPLNLCPICLHKLQCAVGFSIVERYKA LVRWIDDESSDTPGATPEHSHEDNGNLPKPV EAFKEWKEWIIKCLAVLQK
892	2242	A	7723	2	1650	SAPTAPARPCRAERGSGGGMLALLAASVALA VAAGAQDSPAPGSRFVCTALPPEAVHAGCPL PAMPMQGGAQSPEEELRAAVLQLRETVVQQ KETLASARAIRELTGKLARCEGLAGGKARGA GATGKDTMGDLPRDPGHVVEQLSRSLQTLK DRLESLEPLPAMPMQGGAQSPEEELRAAVLQ LRETVVQQKETLASARAIRELTGKLARCEGL AGGKARGAGATGKDTMGDLPRDPGHVVEQ LSRSLQTLKDRLESLEHQLRANVSNAGLPGD FREVLQQRLGELERQLLRKGAELEDEKSLLH NETSAHRQKTESTLNALLQRVTELERGNSAF KSPNAFKVSLPLRTNYLYGKIKKTLPELYAFT ICLWLRSSASPGMGTPFSYAVPGQANEIVLIE WGNNPIELLINDKVAQLPLFVSDGKWHHICV TWTTRDGMWEAFQDGKKLGTGENLAPWHPI KPGGVLILGQEQDTVGGRFDATQAFVGELSQ FNIWDRVLRAQEIVNIANCSTNMPGNIIPWVD NNVDVFGGASKWPVETCEERLLDL
893	2243	Α	7729	3554	2419	LTAGTAMNYPLTLEMDLENLEDLFWELDRL DNYNDTSLVENHLCPATEGPLMASFKAVFVP VAYSLIFLLGVIGNVLVLVILERHRQTRSSTET FLFHLAVADLLLVFILPFAVAEGSVGWVLGTF LCKTVIALHKVNFYCSSLLLACIAVDRYLAIV HAVHAYRHRRLLSIHITCGTIWLVGFLLALPEI LFAKVSQGHHNNSLPRCTFSQENQAETHAWF TSRFLYHVAGFLLPMLVMGWCYVGVVHRLR QAQRRPQRQKAVRVAILVTSIFFLCWSPYHIV IFLDTLARLKAVDNTCKLNGSLPVAITMCEFL GLAHCCLNPMLYTFAGVKFRSDLSRLLTKLG CTGPASLCQLFPSWRRSSLSESENATSLTTF
894	2244	Α	7738	670	287	FVTRAGRWGAGARVRGGAGGMASGAARWL VLAPVRSGALRSGPSLRKDGDVSAAWSGSGR SLVPSRSVIVTRSGAILPKPVKMSFGLLRVFSI VIPFLYVGTLISKNFAALLEEHDIFVPEDDDDD D
895	2245	A	7753	119	278	APYAHSQVHCLDKVCGLLPFLNPEVPDQFYR
						LWLSLFLHAGKEAPHCPRTRPL
896	2246	Α	7754	1	372	SPAWWNSQQRVVSPFLALLTLEPTFHHLLPIM

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897	2247	A	7761	1725	445	TKRGRQVCADPSEEWVQKYVSDLELSA RPRRRGTHHFSCVLGSFRVSAMFPRVSTFLPL RPLSRHPLSSGSPETSAAAIMLLTVRHGTVRY RSSALLARTKNNIQRYFGTNSVICSKKDKQSV RTEETSKETSESQDSEKENTKKDLLGIIKGMK VELSTVNVRTTKPPKRRPLKSLEATLGRLRRA TEYAPKKRIEPLSPELVAAASAVADSLPFDKQ TTKSELLSQLQQHEEESRAQRDAKRPKISFSNI ISDMKVARSATARVRSRPELRIQFDEGYDNYP GQEKTDDLKKRKNIFTGKRLNIFDMMAVTKE APETDTSPSLWDVEFAKQLATVNEQPLQNGF EELIQWTKEGKLWEFPINNEAGFDDDGSEFH EHIFLEKHLESFPKQGPIRHFMELVTCGLSKNP YLSVKQKVEHIEWFRNYFNEKKDILKESNIQF KLRPWKFLFRNN
898	2248	A	7775 *	85	496	SCQTTQPPAQSCSTGTMRIMLLFTAILAFSLA QSFGAVCKEPQEEVVPGGGRSKRDPDLYQLL QRLFKSHSSLEGLLKALSQASTDPKESTSPEK RDMHDFFVGLMGKRSVQPDSPTDVNQENVP SFGILKYPPRAE
899	2249	A	7785	179	703	PFHLGASSNTFRLQVQTQESKAQKEVKMGFI FSKSMNESMKNQKEFMLMNARLQLERQLIM QSEMRERQMAMQIAWSREFLKYFGTFFGLA AISLTAGAIKKKKPAFLVPIVPLSFILTYQYDL GYGTLLERMKGEAEDILETEKSKLQLPRGMIT FESIEKARKEQSRFFIDK
900	2250	A	7789	1465	300	VWLPLKSYKIRSPSLHCQCEIFREEFLFSSLQE GRDKDTFSKMAMVSEFLKQAWFIENEEQEY VQTVKSSKGGPGSAVSPYPTFNPSSDVAALH KAIMVKGVDEATIIDILTKRNNAQRQQIKAAY LQETGKPLDETLKKALTGHLEEVVLALLKTP AQFDADELRAAMKGLGTDEDTLIEILASRTN KEIRDINRVYREELKRDLAKDITSDTSGDFRN ALLSLAKGDRSEDFGVNEDLADSDARALYEA GERRKGTDVNVFNTILTTRSYPQLRRVFQKY TKYSKHDMNKVLDLELKGDIEKCLTAIVKCA TSKPAFFAEKLHQAMKGVGTRHKALIRIMVS RSEIDMNDIKAFYQKMYGISLCQAILDETKGD YEKILVALCGGN
901	2251	A	7796		807	VEFHPQRARAGARAPSMGVLLTQRTLLSLVL ALLFPSMASMAAIGSCSKEYRVLLGQLQKQT DLMQDTSRLLDPYIRIQGLDVPKLREHCRERP GAFPSEETLRGLGRRCFLQTLNATLGCVLHRL ADLEQRLPKAQDLERSGLNIEDLEKLQMARP NILGLRNNIYCMAQLLDNSDTAEPTKAGRGA SQPPTPTPASDAFQRKLEGCRFLHGYHRFMH SVGRVFSKWGESPNRSRRHSPHQALRKGVRR TRPSRKGKRLMTRGQLPR
902	2252	A	7802	2	721	TAARRRQKGTAARRLQKGTAARRRQKGTAA RRRQKGTAARRPQKGTAARRRQKGTAARRR QKGTAARRRQKGTAARRRQKG TAARRRQKGTAARRRQKG TAARRRQKGTAARRRQKGLAIASRGCPCASR AGGVRGAGSRLRAMAPKVFRQYWDIPDGTD CHRKAYSTTSIASVAGLTAAAYRVTLNPPGTF LEGVAKVGQYTFTAAAVGAVFGLTTCISAHV REKPDDPLNYFLGGCAGGLTLGARTHNYGIG AAACVYFGIAASLVKMGRLEGWEVFAKPKV

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
903	2253	Α	7807	1	584	PWLPWSDGRAARSSRKCPRSRFPVQVGKMA VSTVFSTSSLMLALSRHSLLSPLLSVTSFRRFY RGDSPTDSQKDMIEIPLPPWQERTDESIETKR ARLLYESRKRGMLENCILLSLFAKEHLQHMT EKQLNLYDRLINEPSNDWDIYYWATEAKPAP EIFENEVMALLRDFAKNKNKEQRLRAPDLEY LFEKPR
904	2254	A	7813	40	821	GAGRALGHLETGAGDVAAALPARKFPRSLLG AGARLTGWTMNVFRILGDLSHLLAMILLLGK IWRSKCCKGISGKSQILFALVFTTRYLDLFTNF ISIYNTVMKVVFLLCAYVTVYMIYGKFRKTF DSENDTFRLEFLLVPVIGLSFLENYSFTLLEIL WTFSIYLESVAILPQLFMISKTGEAETITHYL FFLGLYRALYLANWIRRYQTENFYDQIAVVS GVVQTIFYCDFFYLYVTKGRSWDDSNADTGL RSYSSI
905	2255	A	7817	1399	881	LSNKDVLSPQLKDENSKLRRKLNEVQSFSEA QTEMVRTLERKLEAKMIKEESDYHDLESVVQ QVEQNLELMTKRAVKAENHVVKLKQEISLL QAQVSNFQRENEALRCGQGASLTVVKQNAD VALQNLRVVMNSAQASIEQLVSGAETLNLVA EILKSIDRISEVKDEEEDS
906	2256	A	7822	3	1462	DSPRNRFEILGRPTRTPTRPGPRPAMEDLDAL LSDLETTTSHMPRSGAPKERPAEPLTPPPSYG HQPQTGSGESSGASGDKDHLYSTVCKPRSPK PAAPAAPPFSSSSGVLGTGLCELDRLLQELNA TQFNITDEIMSQFPSSKVASGEQKEDQSEDKK RPSLPSSPSPGLPKASATSATLELDRLMASLSD FRVQNHLPASGPTQPPVVSSTNEGSPSPPEPTG KGSLDTMLGLLQSDLSRRGVPTQAKGLCGSC NKPIAGQVVTALGRAWHPEHFVCGGCSTAL GGSSFFEKDGAPFCPECYFERFSPRCGFCNQPI RHKMVTALGTHWHPEHFCCVSCGEPFGDEG FHEREGRPYCRRDFLQLFAPRCQGCQGPILDN YISALSALWHPDCFVCRECFAPFSGGSFFEHE GRPLCENHFHARRGSLCATCGLPVTGRCVSA LGRRFHPDHFTCTFCLRPLTKGSFQERAGKPY CQPCFLKLFG
907	2257	A	7828	1792	1671	FIYVNQSFAPSPDQEVGTLYECFGSDGKLVLH YCKSQAWG
908	2258	A	7842	110	1172	KLSCPCSHGTRVTAVRGPRLKAGVQWHDLG SLQPPPSGLKQSSHLSLSSSWDFRHAPTHPET YTCPKMIEMEQAEAQLAELDLLASMFPGENE LIVNDQLAVAELKDCIEKKTMEGRSSKVYFTI NMNLDVSDEKMAMFSLACILPFKYPAVLPEI TVRSVLLSRSQQTQLNTDLTAFLQKHCHGDV CILNATEWVREHASGYVSRDTSSSPTTGSTVQ SVDLIFTRLWIYSHHIYNKCKRKNILEWAKEL SLSGFSMPGKPGVVCVEGPQSACEEFWARLR KLNWKRILIRHREDIPFDGTNDETERQRKFSIF EEKVFSVNGARGNHMDFGQLYQFLNTKGCG DVFQMFLWV
909	2259	A	7870	3067	2923	EGICVYTFIYVHMYTRTCMHTYPYMYMNSV LISSEILLIPSKYLFESK
910	2260	A	7884	212	4874	GALTWSHPLLAVCPQGVWLGSTPSGSPALLP PSHRVNAEPGCVVTNACASGPCPPHANCRDL WQTFSCTCQPGYYGPGCVDACLLNPCQNQG SCRHLPGAPHGYTCDCVGGYFGHHCEHRMD QQCPRGWWGSPTCGPCNCDVHKGFDPNCNK

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SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence		ł	914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
				amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
				peptide		/=possible nucleotide deletion, \=possible nucleotide insertion
				sequence		TNGOCHCKEFHYRPRGSDSCLPCDCYPVGST
						SRSCAPHSQCPCRPGALGRQCNSCDSPFAEV
						TASGCRVLYDACPKSLRSGVWWPOTKFGVL
1 .		ļ				ATVPCPRGALGLRGAGAAVRLCDEAQGWLE
						PDLFNCTSPAFRELSLLLDGLELNKTALDTME
						AKKLAQRLREVTGHTDHYFSQDVRVTARLL
						AHLLAFESHQQGFGLTATQDAHFNENLLWA
						GSALLAPETGDLWAALGQRAPGGSPGSAGLV
1		İ				RHLEEYAATLARNMELTYLNPMGLVTPNIML
1						SIDRMEHPSSPRGARRYPRYHSNLFRGQDAW
						DPHTHVLLPSQSPRPSPSEVLPTSSSIENSTTSS
						VVPPPAPPEPEPGISIIILLVYRTLGGLLPAQFQ
1						AERRGARLPQNPVMNSPVVSVAVFHGRNFLR
						GILESPISLEFRLLQTANRSKAICVQWDPPGLA
1						EQHGVWTARDCELVHRNGSHARCRCSRTGT
		ļ				FGVLMDASPRERLEGDLELLAVFTHVVVAVS
						VAALVLTAAILLSLRSLKSNVRGIHANVAAA
						LGVAELLFLLGIHRTHNQLVCTAVVILLHYFF
					,	LSTFAWLFVQGLHLYRMQVEPRNVDRGAMR
						FYHALGWGVPAVLLGLAVGLDPEGYGNPDF
						CWISVHEPLIWSFAGPVVLVIVMNGTMFLLA
						ARTSCSTGQREAKKTSALTLRSSFLLLLLVSA
						SWLFGLLAVNHSILAFHYLHAGLCGLQGLAV
						LLLFCVLNADARAAWMPACLGRKAAPEEAR PAPGLGPGAYNNTALFEESGLIRITLGASTVSS
1						VSSARSGRTQDQDSQRGRSYLRDNVLVRHGS
						AADHTDHSLQAHAGPTDLDVAMFHRDAGA
	,					DSDSDSDLSLEEERSLSIPSSESEDNGRTRGRF
					'	QRPLCRAAQSERLLTHPKDVDGNDLLSYWPA
						LGECEAAPCALQTWGSERRLGLDTSKDAAN
[						NNQPDPALTSGDETSLGRAQRQRKGILKNRL
						OYPLVPOTRGAPELSWCRAATLGHRAVPAAS
						YGRIYAGGGTGSLSQPASRYSSREQLDLLLRR
1 !			ľ			QLSRERLEEAPAPVLRPLSRPGSQECMDAAPG
						RLEPKDRGSTLPRRQPPRDYPGAMAGRFGSR
1					,	DALDLGAPREWLSTLPPPRRTRDLDPQPPPLP
						LSPQRQLSRDPLLPSRPLDSLSRSSNSREQLDQ
						VPSRHPSREALGPLPQLLRAREDSVSGPSHGP
						STEQLDILSSILASFNSSALSSVQSSSTPLGPHT
						TATPSATASVLGPSTPRSATSHSISELSPDSEPR
						DTQALLSATQAMDLRRRDYHMERPLLNQEH
						LEELGRWGSAPRTHQWRTWLQCSRARAYAL
						LLQHLPVLVWLPRYPVRDWLLGDLLSGLSVA
						IMQLPQGLAYALLAGLPPVFGLYSSFYPVFIY
011	0061		7000	21	906	FLFGTSRHISVESLCVPGPVDT
911	2261	A	7890	21	806	EFGTSRSSRSMAEDLGLSFGETASVEMLPEHG
[ ]						SCRPKARSSSARWALTCCLVLLPFLAGLTTYL
						LVSQLRAQGEACVQFQALKGQEFAPSHQQV YAPLRADGDKPRAHLTVVRQTPTQHFKNQFP
						ALHWEHELGLAFTKNRMNYTNKFLLIPESGD
						YFIYSQVTFRGMTSECSEIRQAGRPNKPDSITV VITKVTDSYPEPTQLLMGTKSVCEVGSNWFQ
						PIYLGAMFSLQEGDKLMVNVSDISLVDYTKE
						DKTFFGAFLL
912	2262	A	7891	1263	111	ACGIRHEGALPGLTATPEAMLRFLPDLAFSFL
714	2202	Λ.	7071	1200	1 1 1	LILALGQAVQFQEYVFLQFLGLDKAPSPQKFQ
						PVPYILKKIFODREAAATTGVSRDLCYVKELG
						VRGNVLRFLPDQGFFLYPKKISQASSCLQKLL
-						YFNLSAIKEREQLTLAQLGLDLGPNSYYNLGP
					ļ	ELELALFLVQEPHVWGQTTPKPGKMFVLRSV
	<u> </u>	<u> </u>	L			

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  PWPQGAVHFNLLDVAKDWNDNPRKNFGLFL
						EILVKEDRDSGVNFQPEDTCARLRCSLHASLL VVTLNPDQCHPSRKRRAAIPVPKLSCKNLCH RHQLFINFRDLGWHKWIIAPKGFMANYCHGE CPFSLTISLNSSNYAFMQALMHAVDPEIPQAV CIPTKLSPISMLYQDNNDNVILRHYEDMVVD ECGCG
913	2263	A	7892	15	849	ASRLPRGPGCGADMRPLLGLLLVFAGCTFAL YLLSTRLPRGRRLGSTEEAGGRSLWFPSDLAE LRELSEVLREYRKEHQAYVFLLFCGAYLYKQ GFAIPGSSFLNVLAGALFGPWLGLLLCCVLTS VGATCCYLLSSIFGKQLVVSYFPDKVALLQR KVEENRNSLFFFLLFLRLFPMTPNWFLNLSAPI LNIPIVQFFFSVLIGLIPYNFICVQTGSILSTLTS LDALFSWDTVFKLLAIAMVALIPGTLIKKFSQ KHLQLNETSTANHIHSRKDT
914	2264	A	7893	815	959	KSGWVWWLTPLIPALWEAQTEGSLRPEVKN RLSNITRPFFSKKKKILV
915	2265	A	7909	3	641	HASGPGGLLRRRRGSGANMPVARSWVCRKT YVTPRRPFEKSRLDQELKLIGEYGLRNKREV WRVKFTLAKIRKAARELLTLDEKDPRRLFEG NALLRRLVRIGVLDEGKMKLDYILGLKIEDFL ERRLQTQVFKLGLAKSIHHAHVLIQQCHIRVR EQVVNILFFTVRLDSQKHIDFSLCFPIGVANPS HVKRKNASKGQGGAGARDDEEEE
916	2266	A	7914	3	967	VAHTQWHTCQRLSQLTHRSILKYLLIDTHAC QVLILKHTHASLSLPSCQECFPSSIPSASHMVS HPHPPPSPRWGQTPEGLPAASPCGPGPRSCFS SILPTGDSWGMLACLCTVLWHLPAVPALNRT GDPGPGPSIQKTYDLTRYLEHQLRSLAGTYLN YLGPPFNEPDFNPPRLGAETLPRATVDLEVW RSLNDKLRLTQNYEAYSHLLCYLRGLNRQAA TAELRRSLAHFCTSLQGLLGSIAGVMAALGY PLPQPLPGTEPTWTPGPAHSDFLQKMDDFWL LKELQTWLWRSAKDFNRLKKKMQPPAAAVT LHLGAHGF
917	2267	A	7921	2	1166	RPRRGQGLVQEVQTENVTVAEGGVAEITCRL HQYDGSIVVIQNPARQTLFFNGTRALKDERFQ LEEFSPRRVRIRLSDARLEDEGGYFCQLYTED THHQIATLTVLVAPENPVVEVREQAVEGGEV ELSCLVPRSRPAATLRWYRDRKELKGVSSSQ ENGKVWSVASTVRFRVDRKDDGGIIICEAQN QALPSGHSKQTQYVLDVQYSPTARIHASQAV VREGDTLVLTCAVTGNPRPNQIRWNRGNESL PERAEAVGETLTLPGLVSADNGTYTCEASNK HGHARALYVLVVYGESRLRPTEGGGAPDP GAVVEAQTSVPYAIVGGILALLVFLIICVLVG MVWCSVRQKGSYLTHEASGLDEQGEAREAF LNGSDGHKRKEEFFI
918	2268	A	7938	3	2653	RRRLPPASPPSSSVSSSLSPSAVVMACRWSTK ESPRWRSALLLLFLAGVYGNGALAEHSENVH ISGVSTACGETPEQIRAPSGIITSPGWPSEYPAK INCSWFIRANPGEIITISFQDFDIQGSRRCNLD WLTIETYKNIESYRACGSTIPPPYISSQDHIWIR FHSDDNISRKGFRLAYFSGKSEEPNCACDQFR CGNGKCIPEAWKCNNMDECGDRSDEEICAKE ANPPTAAAFQPCAYNQFQCLSRFTKVYTCLP ESLKCDGNIDCLDLGDEIDCDVPTCGQWLKY FYGTFNSPNYPDFYPPGSNCTWLIDTGDHRK

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						VILRFTDFKLDGTGYGDYVKIYDGLEENPHK LLRVLTAFDSHAPLTVVSSSGQIRVHFCADKV NAARGFNATYQVDGFCLPWEIPCGGNWGCY TEQQRCDGYWHCPNGRDETNCTMCQKEEFP CSRNGVCYPRSDRCNYQNHCPNGSDEKNCFF CQPGNFHCKNNRCVFESWVCDSQDDCGDGS DEENCPVIVPTRVITAAVIGSLICGLLLVIALG CTCKLYSLRMFERRSFETQLSRVEAELLRREA PPSYGQLIAQGLIPPVEDFPVCSPNQASVLENL RLAVRSQLGFTSVRLPMAGRSSNIWNRIFNFA RSRHSGSLALVSADGDEVVPSQSTSREPERNH THRSLFSVESDDTDTENERRDMAGASGGVAA PLPQKVPPTTAVEATVGACASSSTQSTRGGH ADNGRDVTSVEPPSVSPARHQLTSALSRMTQ GLRWVRFTLGRSSSLSQNQSPLRQLDNGVSG REDDDDVEMLIPISDGSSDFDVNDCSRPLLDL ASDQGQGLRQPYNATNPGVRPSNRDGPCERC GIVHTAQIPDTCLEVTLKNETSDDEALLLC
919	2269	A	7951	1674	1839	VVRVTCCPPARSTTERTNAYDEEDCVEMVAS
920	2270	A	7953	47	572	GGWNDVACHTTMYFMCEFDKKNM GGRASWPEQAKEPRREGHTDKQQTEDVLAA GLRCLPHLPAICARRMSPAFRAMDVEPRAKG VLLEPFVHQVGGHSCVLRFNETTLCKPLVPRE HQFYETLPAEMRKFTPQYKGKSQLLEGLPHW RGDVRDRGHGRPWQPSLEPSLPPTLCFPSLSS FSSSWPSAQHLTPSVFNPW
921	2271	A	7957	612	812	RSGRTVVTGIGYSKALQSSNRNTKSLLQNEF MMVYSFRALSFKESTWATFQHGGEATKSRSL SSTO
922	2272	A	7967	1443	1660	ENITEKWKEIWMCRGNKKSCCWTFIKDRHLT VSCCKSKSGETLLICIFCSNLVGFFFFGIRGFSN WELVKPN
923		A	7981	1	3023	GSAPRAATAMARARPPPPPSPPPGLLPLLPLL LLPLLLLPAGCRALEETLMDTKWVTSELAWT SHPESGWEEVSGYDEAMNPIRTYQVCNVRES SQNNWLRTGFIWRRDVQRVYVELKFTVRDC NSIPNIPGSCKETFNLFYYEADSDVASASSPFW MENPYVKVDTIAPDESFSRLDAGRVNTKVRS FGPLSKAGFYLAFQDQGACMSLISVRAFYKK CASTTAGFALFPETLTGAEPTSLVIAPGTCIPN AVEVSVPLKLYCNGDGEWMVPVGACTCATG HEPAAKESQCRPCPPGSYKAKQGEGPCLPCPP NSRTTSPAASICTCHNNFYRADSDSADSACTT VPSPPRGVISNVNETSLILEWSEPRDLGVRDD LLYNVICKKCHGAGGASACSRCDDNVEFVPR QLGLSEPRVHTSHLLAHTRYTFEVQAVNGVS GKSPLPPRYAAVNITTNQAAPSEVPTLRLHSS SGSSLTLSWAPPERPNGVILDYEMKYFEKSEG IASTVTSQMNSVQLDGLRPDARYVVQVRART VAGYGQYSRPAEFETTSERGSGAQQLQEQLP LIVGSATAGLVFVVAVVVIAIVCLRKQRHGS DSEYTEKLQQYIAPGMKVYIDPFTYEDPNEA VREFAKEIDVSCVKIEEVIGAGEFGEVCRGRL KQPGRREVFVAIKTLKVGYTERQRRDFLSEA SIMGQFDHPNIIRLEGVVTKSRPVMILTEFME NCALDSFLRLNDGQFTVIQLVGMLRGIAAGM KYLSEMNYVHRDLAARNILVNSNLVCKVSDF GLSRFLEDDPSDPTYTSSLGGKIPIRWTAPEAI AYRKFTSASDVWSYGIVMWEVMSYGERPY

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  WDMSNQDVINAVEQDYRLPPPMDCPTALHQ LMLDCWVRDRNLRPKFSQIVNTLDKLIRNAA SLKVIASAQSGMSQPLLDRTVPDYTTFTTVGD WLDAIKMGRYKESFVSAGFASFDLVAQMTA EDLLRIGVTLAGHQKKILSSIQDMRLQMNQT LPVQV
924	2274	A	7985	1	503	FRPRTKKATAMYLEHYLDSIENLPCELQRNF QLMRELDQRTEDKKAEIDILAAEYISTVKTLS PDQRVERLQKIQNAYSKCKEYSDDKVQLAM QTYEMVDKHIRRLDADLARFEADLKDKMEG SDFESSGGRGLKKGRGQKEKRGSRGRGRRTS EEDTPKKKKHKGG
925	2275	A	7994	447	589	LPCSFCAQCMSSFERVWLQQSHFHNPRWNSR
926	2276	A	7996	925	582	SPIRCYCQHWPHCVHC  GPCKVCCITLAIMLQCHSFYRKDVQVEHPKS LNPKYSQIENFLSADMALKRKCLLSISDLDFW IWDAQPVGIMQTLQNLKKIPNPGCFWSQAFQI RDTQPILPLGGRYYITIRQ
927	2277	A	7998	2	353	RIQRPLNSRSPNHSLFVKAELTAKQATMKLSV CLLLVTLALCCYQANAEFCPALVSELLDFFFI SEPLFKLSLAKFDAPPEAVAAKLGVKRCTDQ MSLQKRSLIAEVLVKILKKCSV
928	2278	A	8004	130	588	LAPLRCQPGTRTQPRSHPAANDPSAAMSAAG ARGLRATYHRLLDKVELMLPEKLRPLYNHPA GPRTVFFWAPIMKWGLVCAGLADMARPAEK LSTAQSAVLMATGFIWSRYSLVIIPKNWSLFA VNFFVGAAGASQLFRIWRYNQELKAKAHK
929	2279	A .	8007	2	1016	EFARRRVFIAAREMSLLRSLRVFLVARTGSYP AGSLLRQSPQPRHTFYAGPRLSASASSKELLM KLRRKTGYSFVNCKKALETCGGDLKQAEIWL HKEAQKEGWSKAAKLQGRKTKEGLIGLLQE GNTTVLVEVNCETDFVSRNLKFQLLVQQVAL GTMMHCQTLKDQPSAYSKGFLNSSELSGLPA GPDREGSLKDQLALAIGKLGENMILKRAAWV KVPSGFYVGSYVHGAMQSPSLHKLVLGKYG ALVICETSEQKTNLEDVGRRLGQHVVGMAPL SVGSLDDEPGGEAETKMLSQPYLLDPSITLGQ YVOPOGYSVVDFVRFECGEGEEAAETE
930	2280	A	8008	3	1679	NSRVWGPWTEPSAGSLRPMARKQNRNSKEL GLVPLTDDTSHAGPPGPGRALLECDHLRSGV PGGRRRKDWSCSLLVASLAGAFGSSFLYGYN LSVVNAPTPYIKAFYNESWERRHGRPIDPDTL TLLWSVTVSIFAIGGLVGTLIVKMIGKVLGRK HTLLANNGFAISAALLMACSLQAGAFEMLIV GRFIMGIDGGVALSVLPMYLSEISPKEIRGSLG QVTAIFICIGVFTGQLLGLPELLGKESTWPYLF GVIVVPAVVQLLSLPFLPDSPRYLLLEKHNEA RAVKAFQTFLGKADVSQEVEEVLAESRVQRS IRLVSVLELLRAPYVRWQVVTVIVTMACYQL CGLNAIWFYTNSIFGKAGIPPAKIPYVTLSTGG IETLAAVFSGLVIEHLGRRPLLIGGFGLMGLFF GTLTITLTLQDHAPWVPYLSIVGILAIIASFCSG PGGIPFILTGEFFQQSQRPAAFIIAGTVNWLSN FAVGLLFPFIQKSLDTYCFLVFATICITGAIYL YFVLPETKNRTYAEISQAFSKRNKAYPPEEKI DSAVTDGKINGRP
931	2281	A	8009	861	300	AAGAVVSAMPKAKGKTRRQKFGYSVNRKRL NRNARRKAAPRIECSHIRHAWDHAKSVRQNL AEMGLAVDPNRAVPLRKRKVKAMEVDIEER

PKELVRKPYVLNDLAEASLPEKKGMITSK	SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
DIRGSHPSSPORGESGNYGGETARVLRAGK   GGRGOJPECPHPPPPPPPSGCCRGFHO   SLEAKARHPASVREMRGKVKMRALRAPA   STRASSRQPINE     933   2283   A   8012   147   1077   PPPVPPASRSDMAQNLKDLAGRLPAGFRAMGT   RATASSRQPINE   PPVPPASRSDMAQNLKDLAGRLPAGFRAMGT   RATASSRQPINE							PKELVRKPYVLNDLEAEASLPEKKGNTLSRD LIDYVRYMVENHGEDYKAMARDEKNYYQD TPKQIRSKINVYKRFYPAEWQDFLDSLQKRK MEVE
933   2283   A   8012   147   1077   PPVPPASRSDMAONLKDLAGRIPAGPRGMGT   NGIGGVQQDTILAEGLHFRIPWFQVPIPVDIRA RPRISISPTGSKDLQMVNSEVLKSVPANQEL   PSMYQRLGLDYERF VLPSIVNEVLKSVPANGEL   PSMYQRLGLDYERF VLPSIVNEVLKSVPANGEL   PSMYQRLGLDYERF VLPSIVNEVLKSVPANGEL   PSMYQRLGLDYERF VLPSIVNEVLKSVPANGEL   PSMYQRLGLDYERF VLPSIVNEVLKSVVANGE   NASQLITORAQVSLLIRE IERAKDPSLILDD   VAITELSFSREYTAAVEAKQVAQDEAQRAQF   LVEKAKQEQRQKIVQAGGEAQRAQF   LVEKAKQEQRQKIVQAGGEAQAKMLGEAL   SKNPGYIKLRKIRAAQNISKITATSQNRIVTLTA   DNILVIALQDESFTRGSDSLKGKK   SKPESTAQVLVDSAEGGSLAGARELAAQKREQ   RLRKFRELHLMRNEAKKLNIQEVVEDLKRL   KLPANWEAKKARLEWLKEEKKKECAARG   EDYEVVKLLEISABDAERWERKKKRARDFLGF   FSDYAAAQLQYHELTIKOPDMETYERLE   KHGEEFFFTSNSLLHGTHVPSTEEDIRMYIDLE   KQIEKTDRVSKSRRPYNDDADIDYNDRNAKF   NKKAGRFYGKYTAEIKQNLERGTAV   LVSSTVMLITEREAPWNSTAWTISVYFLKFL   QGGGTGSTGMKDSALTILLGIGFSIRHSLSIRL   SQHSSPAPMYSQTFHILU	932	2282	A	8011	412	1	DIRGSHFSSPQRQRSQRVPGKETARVLRAGK QGRGQIPIPCPWPPPPPPPPPGSPGPGCRQFHQ SLEAKARHPASVREMRGKVKMRRALRRAPA
934   2284   A   8023   255   982   SQFSLSQVLVDSAEEGSLAAARELAAQKREQ   RLRKFRELHLMRNEAKKHHQEVVEEDKRL   KLPANWEAKKARLEWELKEEEKKKECAARG   EDVEKVKLLEISAEDAERWERKKKRKAPDLG   FSDYAAQLRQVIRLITKQIKEPMETYERLER   KHGEEFFFTSNSLLHGTHVPSTEEIDRAVIDLE   KQIEKRDKYSRRRPYNDDADIDYINERNAKF   NKKAERFYGKYTAEIKQNLERGTAV     935   2285   A   8027   59   310   LVSSTVNILITEKAPWNSLAWTVTSYVPLKFL   QGGTGSTGMRDSALTILLGIGPSRHISLSIRL   SQHSSPAPMYSQTFHILVLG     936   2286   A   8032   1   639   SGRECNMAKTYDYLFKLLIGDSGVGKTCVL   FRESDAFNSTFSTIGIDFKIRTIELDGKRIKLQ   INDITAGGERFRITITATYRGAMGIMLVYDIT   NIEKSFDNIRNIKRNEEHASADVEKMILGNKC   DVNDKRQVSKERGEKLALDYGIKFMETSAK   ANINVENAFFTLARDIKAKMDKKLEGNSPQG   SNQGVKITDPQGKSFFRCVLL     937   2287   A   8039   393   311   EETIHSENSYLLEKYPISANLTLTIA     938   2288   A   8052   675   -1334   LHPAATSTAWLHVPPGLSMALSWVLTVLSLL   PLLEAQIPLCANLVPYITNATLDRITGKWFYT   ASAFRNEEYNKSVQEIQAFFFYFTPNKTEDTIFT   LREYGTRODQCTYNTTYLNVQRENGTISRYV   GGQEHFA-ILLILRDTKTYMLAFDVNDEKNW   GLSVYADKPETTKEQLGFFYEALDCLRIPKSD   VVYTDWKKDKCEPLERQHEKERKQEEGES     939   2289   A   8055   12   1039   SSVAEFPERVOLSQPONWNFSGAGGAWSLDF   AFQLKWSAELARGESIMDGKGGMDGSKP   AGPRDFPGIRLLSNPLMGDAVSDWSPMHEAA   HIGHQLSLRNLISQGWAVNIITADHYSPLHEA   CLGGHLSCVKILLKHGAQVNGVTADWHTPL   FNACVSGSWDCVNILLLQHGASVQPESDLASP   HEAARRGHVECVNSLLAYGGGIDHKISHLGT   PLYLACENQRACVKKLLESGADVNQGKQQ   DSPLHAVARTASEELACLLMDFGADTQAKN   AEGKRPVELVPPESPLAQHLTREGPPSLMQL   CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH   LREGGRESTARGESTARGER   LREGGRESTARGE   CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH   LREGGRESTARGE   CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH   LREGGRESTARGE   CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH   LREGGRESTARGE   CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH   LREGGRESTARGE   CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH   LREGGRESTARGE   CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH   LREGGRESTARGE   CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH   LREGGRESTARGE   CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH   LREGGRESTARGE   CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH   LREGGRESTARGE   CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH   LREGGRESTARGE   CRLRIRKCFGIQQHIGHT   CRLRIRGE   CRLRIRGT	933	2283	A	8012	147	1077	PPVPPASRSDMAQNLKDLAGRLPAGPRGMGT ALKLLLGAGAVAYGVRESVFTVEGGHRAIFF NRIGGVQQDTILAEGLHFRIPWFQYPIIYDIRA RPRKISSPTGSKDLQMVNISLRVLSRPNAQEL PSMYQRLGLDYEERVLPSIVNEVLKSVVAKF NASQLITQRAQVSLLIRRELTERAKDFSLILDD VAITELSFSREYTAAVEAKQVAQQEAQRAQF LVEKAKQEQRQKIVQAEGEAEAAKMLGEAL SKNPGYIKLRKIRAAQNISKTIATSQNRIYLTA
935   2285   A   8027   59   310	934	2284	A	8023	255 .	982	SQFSLSQVLVDSAEEGSLAAAAELAAQKREQ RLRKFRELHLMRNEARKLNHQEVVEEDKRL KLPANWEAKKARLEWELKEEEKKKECAARG EDYEKVKLLEISAEDAERWERKKKRKNPDLG FSDYAAAQLRQYHRLTKQIKPDMETYERLRE KHGEEFFPTSNSLLHGTHVPSTEEIDRMVIDLE KQIEKRDKYSRRRPYNDDADIDYINERNAKF
936   2286	935	2285	A	8027	59	310	LVSSTVNLLTEKAPWNSLAWTVTSYVFLKFL QGGGTGSTGMRDSALTLLGIGPSHRHSLSIRL
937 2287 A 8039 393 311 EETIHSENSYILEKYIPISANLTLTIA 938 2288 A 8052 675 1334 LHPAATSTAWLHVPPGLSMALSWVLTVLSLL PLEAQIPLCANLVPVPITNATLDRITGKWFYI ASAFRNEEYNKSVQEIQATFFYFPNKTEDTIF LREYQTRQDQCIYNTTYLNVQRENGTISRYV GGQEHFAHLLILRDTKTYMLAFDVNDEKNW GLSVYADKPETTKEQLGEFYEALDCLRIPKSD VVYTDWKKDKCEPLEKQHEKERKQEEGES  939 2289 A 8055 12 1039 SSVAEFPERVQLSQPQNWNFSGAGGAWSLDF AEQLKWSAELARLGESIMDGKQGGMDGSKP AGPRDFPGIRLLSNPLMGDAVSDWSPMHEAA IHGHQLSLRNLISQGWAVNIITADHVSPLHEA CLGGHLSCVKILLKHGAQVNGVTADWHTPL FNACVSGSWDCVNLLLQHGASVQPESDLASP IHEAARRGHVECVNSLIAYGGNIDHKISHILGT PLYLACENQQRACVKKLLESGADVNQGKGQ DSPLHAVARTASEELACLLMDFGADTQAKN AEGKRPVELVPPESPLAQLFLEREGPPSLMQL CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH L  940 2290 A 8058 2 1203 KVLSIREPAHSTARKASEPSQPSQPSQPSGPHLI			A	•		639	FRFSEDAFNSTFISTIGIDFKIRTIELDGKRIKLQ IWDTAGQERFRTITTAYYRGAMGIMLVYDIT NEKSFDNIRNWIRNIEEHASADVEKMILGNKC DVNDKRQVSKERGEKLALDYGIKFMETSAK ANINVENAFFTLARDIKAKMDKKLEGNSPQG
938 2288 A 8052 675 1334 LHPAATSTAWLHVPPGLSMALSWVLTVLSLL PLLEAQIPLCANLVPVPITNATLDRITGKWFYI ASAFRNEEYNKSVQEIQATFFYFTPNKTEDTIF LREYQTRQDQCIYNTTYLNVQRENGTISRYV GGQEHFAHLLILRDTKTYMLAFDVNDEKNW GLSVYADKPETTKEQLGEFYEALDCLRIPKSD VVYTDWKKDKCEPLEKQHEKERKQEEGES  939 2289 A 8055 12 1039 SSVAEFPERVQLSQPQNWNFSGAGGAWSLDF AEQLKWSAELARLGESIMDGKQGGMDGSKP AGPRDFPGIRLLSNPLMGDAVSDWSPMHEAA IHGHQLSLRNLISQGWAVNIITADHVSPLHEA CLGGHLSCVKILLKHGAQVNGVTADWHTPL FNACVSGSWDCVNLLLQHGASVQPESDLASP IHEAARRGHVECVNSLIAYGGNIDHKISHLGT PLYLACENQQRACVKKLLESGADVNQGKGQ DSPLHAVARTASEELACLLMDFGADTQAKN AEGKRPVELVPPESPLAQLFLEREGPPSLMQL CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH L	937	2287	A	8039	393	311	
939 2289 A 8055 12 1039 SSVAEFPERVQLSQPQNWNFSGAGGAWSLDF AEQLKWSAELARLGESIMDGKQGGMDGSKP AGPRDFPGIRLLSNPLMGDAVSDWSPMHEAA IHGHQLSLRNLISQGWAVNIITADHVSPLHEA CLGGHLSCVKILLKHGAQVNGVTADWHTPL FNACVSGSWDCVNLLLQHGASVQPESDLASP IHEAARRGHVECVNSLIAYGGNIDHKISHLGT PLYLACENQQRACVKKLLESGADVNQGKGQ DSPLHAVARTASEELACLLMDFGADTQAKN AEGKRPVELVPPESPLAQLFLEREGPPSLMQL CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH L  940 2290 A 8058 2 1203 KVLSIREPAHSTARKASEPSQPSQPSQPGGHLI							LHPAATSTAWLHVPPGLSMALSWVLTVLSLL PLLEAQIPLCANLVPVPITNATLDRITGKWFYI ASAFRNEEYNKSVQEIQATFFYFTPNKTEDTIF LREYQTRQDQCIYNTTYLNVQRENGTISRYV GGQEHFAHLLILRDTKTYMLAFDVNDEKNW GLSVYADKPETTKEQLGEFYEALDCLRIPKSD
		2289	A		12	1039	SSVAEFPERVQLSQPQNWNFSGAGGAWSLDF AEQLKWSAELARLGESIMDGKQGGMDGSKP AGPRDFPGIRLLSNPLMGDAVSDWSPMHEAA IHGHQLSLRNLISQGWAVNIITADHVSPLHEA CLGGHLSCVKILLKHGAQVNGVTADWHTPL FNACVSGSWDCVNLLLQHGASVQPESDLASP IHEAARRGHVECVNSLIAYGGNIDHKISHLGT PLYLACENQQRACVKKLLESGADVNQGKGQ DSPLHAVARTASEELACLLMDFGADTQAKN AEGKRPVELVPPESPLAQLFLEREGPPSLMQL CRLRIRKCFGIQQHHKITKLVLPEDLKQFLLH
	940	2290	A	8058	2	1203	

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						VVDTVMCPNMPNKSVLLYTLSFIYIFIFVIGMI ANSVVVWVNIQAKTTGYDTHCYILNLAIADL WVVLTIPVWVVSLVQHNQWPMGELTCKVTH LIFSINLFGSIFFLTCMSVDRYLSITYFTNTPSS RKKMVRRVVCILVWLLAFCVSLPDTYYLKT VTSASNNETYCRSFYPEHSIKEWLIGMELVSV VLGFAVPFSIIAVFYFLLARAISASSDQEKHSS RKIIFSYVVVFLVCWLPYHVAVLLDIFSILHYI PFTCRLEHALFTALHVTQCLSLVHCCVNPVL YSFINRNYRYELMKAFIFKYSAKTGLTKLIDA SRVSETEYSALEQSTK
941	2291	A	8059	73	432	DMAGLMTIVTSLLFLGVCAHHIIPTGSVVLPS PCCMFFVSKRIPENRVVSYQLSSRSTCLKAGV IFTTKKGQQFCGDPKQEWVQRYMKNLDAKQ KKASPRARAVAVKGPVQRYPGNQTTC
942	2292	A	8067	278	1262	GGIGEIKQRPSCLGRCLDPSLSVLMNISLGLGS VFSAVISQKPSRDICQRGTSLTIQCQVDSQVT MMFWYRQQPGQSLTLIATANQGSEATYESGF VIDKFPISRPNLTFSTLTVSNMSPEDSSIYLCSA GRQGTYEQYFGPGTRLTVTEDLKNVFPPEVA VFEPSEAEISHTQKATLVCLATGFYPDHVELS WWVNGKEVHSGVSTDPQPLKEQPALNDSRY CLSSRLRVSATFWQNPRNHFRCQVQFYGLSE NDEWTQDRAKPVTQIVSAEAWGRADCGFTS ESYQQGVLSATILYEILLGKATLYAVLVSALV LMAMVKRKDSRG
943	2293	A	8070	1	879	MVKVVPATRGNLPRSQLTGTHQHCQPREPKI TASERLRRPRATARLRAHAAPPEPPLAVFAP PSDRKELLALPVACDPVIASVMSWVQAASLI QGPGDKGDVFDEEADESLLAQREWQSNMQR RVKEGYRDGIDAGKAVTLQQGFNQGYKKGA EVILNYGRLRGTLSALLSWCHLHNNNSTLINK INNLLDAVGQCEEYVLKHLKSITPPSHVVDLL DSIEDMDLCHVVPAEKKIDEAKDERLCENNA EFNKNCSKSHSGIDCSYVECCRTQEHAHSGK PKPHMDFGTDSQF
944	2294	A	8073	1	797	ESARWSRQLRRTLIRLSFPISCGRSHAFGGCK MAATSGTDEPVSGELVSVAHALSLPAESYGN DPDIEMAWAMRAMQHAEVYYKLISSVDPQF LKLTKVDDQIYSEFRKNFETLRIDVLDPEELK SESAKEKWRPFCLKFNGIVEDFNYGTLLRLD CSQGYTEENTIFAPRIQFFAIEIARNREGYNKA VYISVQDKEGEKGVNNGGEKRADSGEEENT KNGGEKGADSGEEKEEGINREDKTDKGGEK GKEADKEINKSGEKAM
945	2295	A .	8074	2	505	GAATLLRSASSAARKAAEAEQVWLHLHRYL SADRRVLGLREWGRPASERECSLCQRLKREL NMGDVEKGKKIFIMKCSQCHTVEKGGKHKT GPNLHGLFGRKTGQAPGYSYTAANKNKGIIW GEDTLMEYLENPKKYIPGTKMIFVGIKKKEER ADLIAYLKKATNE
946	2296	A	8081	42	590	EGRRGKFGGKLCNFLFYFHSNSAESRMDVLF VAIFAVPLILGQEYEDEERLGEDEYYQVVYY YTVTPSYDDFSADFTIDYSIFESEDRLNRLDK DITEAIETTISLETARADHPKPVTVKPVTTEPQ SPRSEAMPCPVLRSPIPLPPVRVPLFRWGCISC KKVGRRLLMTLWMGVWQEEIGR
947	2297	A	8084	322	549	GGGSSPRELAGAAGLTVTSQAVAARRQQPSF SRARAPAHSLRAALSLASSARSWGAVSRDRG

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948	2298	В	8093	3905	846	PCPPAIMYQSSNKC  MEPGEVKDRILENISLSVKKLQSYFAACEDEI PAIRNIIDKVLQRLCEHLDHALLYGLQDLSSG YWVLVVHFTRREAIKQIEVLQHVATNLGRSR AWLYLALNENSLESYLRLFQENLGLLHKYYV KNALVCSHDHLTLFLTLVSGLEFIRFELDLDA PYLDLAPYMPDYYKPQYLLDFEDRLPSSVHG SDSLSLNSFNSVTSTNLEWDDSAIAPSSEDYD FGDVFPAVPSVPSTDWEDGDLTDTVSGPRST ASDLTSSKASTRSPTQRQNPFNEEPAETVSSS DTTPVHTTSQEKEEAQALDPPDACTELEVIRV TKKKKIGKKKKSRSDEEASPLHPACSQKKCA KQGDGDSRNGSPSLGRDSPDTMLASPQEEGE GPSSTTESSERSEPGLLIPEMKDTSMERLGQPL SKVIDQLNGQLDPSTWCSRAEPPDQSFRTGSP GDAPERPPLCDFSEGLSAPMDFYRFTVESPST VTSGGGHHDPAGLGQPLHVPSSPEAAGQEEE GGGGEGQTPRPLEDTTREAQELEAQLSLVRE GPVSEPEPGTQEVLCQLKRDQPSPCLSSAEDS GVDEGQGSPSEMVHSSEFRVDNNHLLLLMIH VFRENEEQLFKMIRMSTGHMEGNLQLLYVLL TDCYVYLLRKGATEKPYLVEEAVSYNELDY VSVGLDQQTVKLVCTNRRKQFLLDTADVAL AEFFLASLKSAMIKGCREPPYPSILTDATMEK LALAKFVAQESKCEASAVTVRFYGLVHWED PTDESLGPTPCHCSPPEGTITKEGMLHYKAGT SYLGKEHWKTCFVVLSNGILYQYPDRTDVIP LLSVNMGGEQCGGCRRANTTDRPHAFQVILS DPPCLELSAESEAEMAEWMQHLCQAVSKGVI PQGVAPSPCIPCCLVLTDDRLFTCHEDCQTSF FRSLGTAKLGDISAVSTEPGKEYCVLEFSQDS QQLLPPWVIYLSCTSELDRLLSALNSGWKTIY QVDLPHTAIQEASNKKKFEDALSLIHSAWQR SDSLCRGRASRDPWC*
949	2299	A	8095	9	2374	ARRADTVLLESPSMLQGLLPVSLLLSVAVSAI KELPGVKKYEVVYPIRLHPLHKREAKEPEQQ EQFETELKYKMTINGKIAVLYLKKNKNLLAP GYTETYYNSTGKEITTSPQIMDDCYYQGHILN EKVSDASISTCRGLRGYFSQGDQRYFIEPLSPI HRDGQEHALFKYNPDEKNYDSTCGMDGVL WAHDLQQNIALPATKLVKLKDRKVQEHEKY IEYYLVLDNGEFKRYNENQDEIRKRVFEMAN YVNMLYKKLNTHVALVGMEIWTDKDKIKIT PNASFTLENFSKWRGSVLSRRKRHDIAQLITA TELAGTTVGLAFMSTMCSPYSVGVVQDHSD NLLRVAGTMAHEMGHNFGMFHDDYSCKCPS TICVMDKALSFYIPTDFSSCSRLSYDKFFEDKL SNCLFNAPLPTDIISTPICGNQLVEMGEDCDC GTSEECTNICCDAKTCKIKATFQCALGECCEK CQFKKAGMVCRPAKDECDLPEMCNGKSGNC PDDRFQVNGFPCHHGKGHCLMGTCPTLQEQ CTELWGPGTEVADKSCYNRNEGGSKYGYCR RVDDTLIPCKANDTMCGKLFCQGGSDNLPW KGRIVTFLTCKTFDPEDTSQEIGMVANGTKCG DNKVCINAECVDIEKAYKSTNCSSKCKGHAV CDHELQCQCEEGWIPPDCDDSSVVFHFSIVVG VLFPMAVIFVVVAMVIRHQSSREKQKKDQRP LSTTGTRPHKQKRKPQMVKAVQPQEMSQMK PHVYDLPVEGNEPPASFHKDTNALPPTVFKD

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950	2300	A	8100	1	1251	MGLLLMILASAVLGSFLTLLAQFFLLYRRQPE PPADEAARAGEGFRYIKPVPGLLLREYLYGG GRDEEPSGAAPEGGATPTAAPETPAPPTRETC YFLNATILFLFRELRDTALTRRWVTKKIKVEF EELLQTKTAGRLLEGLSLRDVFLGETVPFIKTI RLVRPVVPSATGEPDGPEGEALPAACPEELAF EAEVEYNGGFHLAIDVDLVFGKSAYLFVKLS RVVGRLRLVFTRVPFTHWFFSFVEDPLIDFEV RSQFEGRPMPQLTSIIVNQLKKIKRKHTLPNY KIRFKPFFPYQTLQGFEEDEEHIHIQWALTE GRLKVTLLECSRLLIFGSYDREANVHCTLELS SSVWEEKQRSSIKTGTISLTAVFMGWHRVSE AFPGLWYKLLVDLPFWGLEDGGPLLTVPLRQ CPG
951	2301	A	8108	1612	839	EVALFCFEMAAGMYLEHYLDSIENLPFELQR NFQLMRDLDQRTEDLKAEIDKLATEYMSSAR SLSSEEKLALLKQIQEAYGKCKEFGDDKVQL AMQTYEMVDKHIRRLDTDLARFEADLKEKQI ESSDYDSSSSKGKKKGRTQKEKKAARARSKG KNSDEEAPKTAQKKLKLVRTSPEYGMPSVTF GSVHPSDVLDMPVDPNEPTYCLCHQVSYGE MIGCDNPDCSIEWFHFACVGLTTKPRGKWFC PRCSQERKKK
952	2302	A	8112	595	291	PSVASLARRFSGRALWPPSHSVPGNRALCPRL LHGTTLPGGNQRELARQKNMKKQSDSVKGK RRDDGLSAAARKQRDSTPRDSEIMQQKQKK ANEKKEEPK
953	2303	A	8118	1	669	VCAGIRDPCSTPLAKPAAGGAENLSFGKQPG LETNILKMTTPNKTPPGADPKQLERTGTVREI GSQAVWSLSSCKPGFGVDQLRDDNLETYWQ SDGSQPHLVNIQFRRKTTVKTLCIYADYKSDE SYTPSKISVRVGNNFHNLQEIRQLELVEPSGW IHVPLTDNHKKPTRTFMIQIAVLANHQNGRD THMRQIKIYTPVEESSIGKFPRCTTIDFMMYRS IR
954	2304	A	8133	66	1015	PPLPPRSFPNLFSRPEPLPEPGRRGCNRSREPA ARAPSPPPPFEGAPGRAMVKVTFNSALAQKE AKKDEPKSGEEALIIPPDAVAVDCKDPDDVV PVGQRRAWCWCMCFGLAFMLAGVILGGAY LYKYFALQPDDVYYCGIKYIKDDVILNEPSAD APAALYQTIEENIKIFEEEEVEFISVPVPEFADS DPANIVHDFNKKLTAYLDLNLDKCYVIPLNT SIVMPPRNLLELLINIKAGTYLPQSYLIHEHMV ITDRIENIDHLGFFIYRLCHDKETYKLQRRETI KGIQKREASNCFAIRHFENKFAVETLICS
955	2305	A	8143	35	1171	VESRSAWHEGEDQIDRLDFIRNQMNLLTLDV KKKIKEVTEEVANKVSCAMTDEICRLSVLVD EFCSEFHPNPDVLKIYKSELNKHIEDGMGRNL ADRCTDEVNALVLQTQQEIIENLKPLLPAGIQ DKLHTLIPCKKFDLSYNLNYHKLCSDFQEDIV FRFSLGWSSLVHRFLGPRNAQRVLLGLSEPIF QLPRSLASTPTAPTTPATPDNASQEELMITLVT GLASVTSRTSMGIIIVGGVIWKTIGWKLLSVS LTMYGALYLYERLSWTTHAKERAFKQQFVN YATEKLRMIVSSTSANCSHQVKQQIATTFARL CQQVDITQKQLEEEIARLPKEIDQLEKIQNNS KLLRNKAVQLENELENFTKQFLPSSNEES
956	2306	A	8157	1854	798	ASGSPAPSSSSAMAAACGPGAAGYCLLLGLH LFLLTAGPALGWNDPDRMLLRDVKALTLHY

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycinc, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  DRYTTSRRLDPIPQLKCVGGTAGCDSYTPKVI
0.55						QCQNKGWDGYDVQWECKTDLDIAYKFGKT VVSCEGYESSEDQYVLRGSCGLEYNLDYTEL GLQKLKESGKQHGFASFSDYYYKWSSADSC NMSGLITIVVLLGIAFVVYKLFLSDGQYSPPP YSEYPPFSHRYQRFTNSAGPPPPGFKSEFTGPQ NTGHGATSGFGSAFTGQQGYENSGPGFWTGL GTGGILGYLFGSNRAATPFSDSWYYPSYPPSY PGTWNRAYSPLHGGSGSYSVCSNSDTKTRTA SGYGGTRRR
957	2307	Α	8159	1492	528	THVVMTGMCYAPHQVLSYINGVTTSKPGVSL VYSMPSRNLSLRLEGLQEKDSGPYSCSVNVQ DKQGKSRGHSIKTLELNVLVPPAPPSCRLQGV PHVGANVTLSCQSPRSKPAVQYQWDRQLPSF QTFFAPALDVIRGSLSLTNLSSSMAGVYVCKA HNEVGTAQCNVTLEVSTGPGAAVVAGAVVG TLVGLGLLAGLVLLYHRRGKALEEPANDIKE DAIAPRTLPWPKSSDTISKNGTLSSVTSARAL RPPHGPPRPGALTPTPSLSSQALPSPRLPTTDG AHPQPISPIPGGVSSSGLSRMGAVPVMVPAQS QAGSLV
958	2308	Α	8161	2340	1192	ELARRPKQQSSEKSRNMIRNWLTIFILFPLKLV EKCESSVSLTVPPVVKLENGSSTNVSLTLRPP LNATLVITFEITFRSKNITILELPDEVVVPPGVT NSSFQVTSQNVGQLTVYLHGNHSNQTGPRIR FLVIRSSAISIINQVIGWIYFVAWSISFYPQVIM NWRRKSVIGLSFDFVALNLTGFVAYSVFNIGL LWVPYIKEQFLLKYPNGVNPVNSNDVFFSLH AVVLTLIIIVQCCLYERGGQRVSWPAIGFLVL AWLFAFVTMIVAAVGVITWLQFLFCFSYIKL AVTLVKYFPQAYMNFYYKSTEGWSIGNVLL DFTGGSFSLLQMFLQSYNNDQWTLIFGDPTK FGLGVFSIVFDVVFFIQHFCLYRKRPGYDQLN
959	2309	A	8163	521	1345	GERAGRRRGRLGVWAQPQPLLPRPVGSRRE MQPPGPPPAYAPTNGDFTFVSSADAEDLSGSI ASPDVKLNLGGDFIKESTATTFLRQRGYGWL LEVEDDDPEDNKPLLEELDIDLKDIYYKIRCV LMPMPSLGFNRQVVRDNPDFWGPLAVVLFFS MISLYGQFRVVSWIITIWIFGSLTIFLLARVLG GEVAYGQVLGVIGYSLLPLIVIAPVLLVVGSF EVVSTLIKLFGVFWAAYSAASLLVGEEFKTK KPLLIYPIFLLYIYFLSLYTGV
960	2310	A	8167	1	2921	MTCFKGQKGEQRSHAFEANKDHKAKVPSPN LYSQLNALQFTVDERSILWLNQFLLDLKQSL NQFMAVYKLNDNSKSDEHVDVRVDGLMLK FVIPSEVKSECHQDQPRAISIQSSEMIATNTRH CPNCRHSDLEALFQDFKDCDFFSKTYTSFPKS CDNFNLLHPIFQRHAHEQDTKMHEIYKGNITP QLNKNTLKTSAATDVWAVYFSQFWIDYEGM KSGKGRPISFVDSFPLSIWICQPTRYAESQKEP QTCNQVSLNTSQSESSDLAGRLKRKKLLKEY YSTESEPLTNGGQKPSSSDTFFRFSPSSSEADI HLLVHVHKHVSMQINHYQYLLLLFLHESLILL SENLRKDVEAVTGSPASQTSICIGILLRSAELA LLLHPVDQANTLKSPVSESVSPVVPDYLPTEN GDFLSSKRKQISRDINRIRSVTVNHMSDNRSM SVDLSHIPLKDPLLFKSASDTNLQKGISFMDY LSDKHLGKISEDESSGLVYKSGSGEIGSETSD KKDSFYTDSSSVLNYREDSNILSFDSDGNQNI LSSTLTSKGNETIESIFKAEDLLPEAASLSENL

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  DISKEETPPVRTLKSQSSLSGKPKERCPPNLAP LCVSYKNMKRSSSQMSLDTISLDSMILEEQLL ESDGSDSHMFLEKGNKKNSTTNYRGTAESVN AGANLQNYGETSPDAISTNSEGAQENHDDLM SVVVFKITGVNGEIDIRGEDTEICLQVNQVTP DQLGNISLRHYLCNRPVGSDQKAVIHSKSSPE
						ISLRFESGPGAVIHSLLAEKNGFLQCHIENFST EFLTSSLMNIQHFLEDETVATVMPMKIQVSNT KINLKDDSPRSSTVSLEPAPVTVHIDHLVVER SDDGSFHIRDSHMLNTGNDLKENVKSDSVLL TSGKYDLKKQRSVTQATQTSPGVPWPSQSAN FPEFSFDFTREQLMEENESLKQELAKAKMAL AEAHLEKDALLHHIKKMTVE
961	2311	A	8172	1442	682	TAAMSIFTPTNQIRLTNVAVVRMKRAGKRFEI ACYKNKVVGWRSGVEKDLDEVLQTHSVFVN VSKGQVAKKEDLISAFGTDDQTEICKQILTKG EVQVSDKERHTQLEQMFRDIATIVADKCVNP ETKRPYTVILIERAMKDIHYSVKTNKSTKQQA LEVIKQLKEKMKIERAHMRLRFILPVNEGKKL KEKLKPLIKVIESEDYGQQLEIVCLIDPGCFREI DELIKKETKGKGSLEVLNLKDVEEGDEKFE
962	2312	A	8175	286	587	NISNKAEVSSHPSVISHSMDSFGQPRPEDNQS VLRRMQKKYWKTKQVFIKATGKKEDEHLVA SDAELDAKLEVFHSVQETCTELLKIIEKYQLR LNGMKS
963	2313	Ą	8181	13	2215	AEGCAERRGTEPVVELSMSWESGAGPGLGSQ GMDLVWSAWYGKCVKGKGSLPLSAHGIVV AWLSRAEWDQVTVYLFCDDHKLQRYALNRI TVWRSRSGNELPLAVASTADLIRCKLLDVTG GLGTDELRLLYGMALVRFVNLISERKTKFAK VPLKCLAQEVNIPDWIVDLRHELTHKKMPHI NDCRRGCYFVLDWLQKTYWCRQLENSLRET WELEEFREGIEEEDQEEDKNIVVDDITEQKPE PQDDGKSTESDVKADGDSKGSEEVDSHCKK ALSHKELYERARELLVSYEEEQFTVLEKFRYL PKAIKAWNNPSPRVECVLAELKGVTCENREA VLDAFLDDGFLVPTFEQLAALQIEYEENVDL NDVLVPKPFSQFWQPLLRGLHSQNFTQALLE RMLSELPALGISGIRPTYILRWTVELIVANTKT GRNARRFSAGQWEARRGWRLFNCSASLDWP RMVESCLGSPCWASPQLLRIIFKAMGQGLPD EEQEKLLRICSIYTQSGENSLVQEGSEASPIGK SPYTLDSLYWSVKPASSSFGSEAKAQQCEQ GSVNDVKEEEKEEKEVLPDQVEEEEENDDQE EEEEDEDDEDDEEEDRMEVGPFSTGQESPTA ENARLLAQKRGALQGSAWQVSSEDVRWDTF PLGRMPGQTEDPAELMLENYDTMYLLDQPV LEQRLEPSTCKTDTLGLSCGVGSGNCSNSSSS
964	2314	A	8184	6	1393	NFEGLLWSQGQLHGLKTGLQLF EPRRNFRDDSTRPRTRGRTRGRRRACRSAE GTGLRSLLLPPRLQLPAGPFSRCRWDPVSSPR PSTMPPKKGGDGIKPPPIIGRFGTSLKIGIVGLP NVGKSTFFNVLTNSQASAENFFFCTIDPNESR VPVPDERFDFLCQYHKPASKIPAFLNVVDIAG LVKGAHNGQGLGNAFLSHISACDGIFHLTRA FEDDDITHVEGSVDPIRDIEIIHEELQLKDEEMI GPIIDKLEKVAVRGGDKKLKPEYDIMCKVKS WVIDQKKPVRFYHDWNDKEIEVLNKHLFLTS KPMVYLVNLSEKDYIRKKNKWLIKIKEWVD KYDPGALVIPFSGALELKLQELSAEERQKYLE

SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq- uence	ĺ	USSN 09/496	location	corresponding to last amino	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline,
seq- uence	uence		914	correspondi	acid residue	Q=Glutamine, R=Arginine, S=Serine,
defice			714	amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
}		]		peptide	Sequence	/=possible nucleotide deletion, \=possible
	1			sequence		nucleotide insertion
		·				ANMTQSALPKIIKAGFAALQLEYFFTAGPDEV
						RAWTIRKGTKAPQAAGKIHTDFEKGFIMAEV
		ľ	Ï		Ì	MKYEDFKEEGSENAVKAAGKYRQQGRNYIV
						EDGDIIFFKFNTPQQPKKK
965	2315	A	8195	1437	594	RSFSLSFSLLSPSEMMALGAAGATRVFVAMV
]		ļ	ļ		ļ	AAALGGHPLLGVSATLNSVLNSNAIKNLPPPL
						GGAAGHPGSAVSAAPGILYPGGNKYQTIDNY
						QPYPCAEDEECGTDEYCASPTRGGDAGVQIC
						LACRKRRKRCMRHAMCCPGNYCKNGICVSS
		ĺ	İ			DQNHFRGEIEETITESFGNDHSTLDGYSRRTT
						LSSKMYHTKGQEGSVCLRSSDCASGLCCARH
						FWSKICKPVLKEGQVCTKHRRKGSHGLEIFQ
0.55	0011	ļ		1	1005	RCYCGEGLSCRIQKDHHQASNSSRLHTCQRH
966	2316	A	8207	416	4082	KFKLIKIMLLTLIILLPVVSKFSFVSLSAPQHW
						SCPEGTLAGNGNSTCVGPAPFLIFSHGNSIFRI
		1			1	DTEGTNYEQLVVDAGVSVIMDFHYNEKRIY
ĺ		ĺ			1	WVDLERQLLQRVFLNGSRQERVCNIEKNVSG
						MAINWINEEVIWSNQQEGIITVTDMKGNNSHI LLSALKYPANVAVDPVERFIFWSSEVAGSLY
		ĺ				RADLDGVGVKALLETSEKITAVSLDVLDKRL
						FWIQYNREGSNSLICSCDYDGGSVHISKHPTO
J			]			HNLFAMSLFGDRIFYSTWKMKTIWIANKHTG
						KDMVRINLHSSFVPLGELKVVHPLAQPKAED
		1				DTWEPEQKLCKLRKGNCSSTVCGQDLQSHLC
						MCAEGYALSRDRKYCEGNDWKYCEDVNEC
1 1		ĺ	ĺ			AFWNHGCTLGCKNTPGSYYCTCPVGFVLLPD
						GKRCHQLVSCPRNVSECSHDCVLTSEGPLCF
						CPEGSVLERDGKTCSGCSSPDNGGCSQLCVPL
1 1			1	1		SPVSWECDCFPGYDLQLDEKSCAASGPQPFL
						LFANSQDIRHMHFDGTDYGTLLSQQMGMVY
						ALDHDPVENKIYFAHTALKWIERANMDGSQ
						RERLIEEGVDVPEGLAVDWIGRRFYWTDRGK
[ [		[		1		SLIGRSDLNGKRSKIITIENISQPRGIAVHPMAK
						RLFWTDTGINPRIESSSLQGLGRLVIASSDLIW
						PSGITIDFLTDKLYWCDAKQSVIEMANLDGSK
1		}	1	l		RRRLTQNDVGHPFAVAVFEDYVWFSDWAMP
						SVIRVNKRTGKDRVRLQGSMLKPSSLVVVHP
						LAKPGADPCLYQNGGCEHICKKRLGTAWCS
						CREGFMKASDGKTCLALDGHQLLAGGEVDL KNQVTPLDILSKTRVSEDNITESQHMLVAEIM
						VSDQDDCAPVGCSMYARCISEGEDATCQCLK
1			1			GFAGDGKLCSDIDECEMGVPVCPPASSKCINT
						EGGYVCRCSEGYQGDGIHCLDIDECQLGVHS
			1			CGENASCTNTEGGYTCMCAGRLSEPGLICPD
			1	1		STPPPHLREDDHHYSVRNSDSECPLSHDGYCL
1			1	1		HDGVCMYIEALDKYACNCVVGYIGERCOYR
			1			DLKWWELRHAGHGQQQKVIVVAVCVVVLV
	ad .		!			MLLLLSLWGAHYYRTQKLLSKNPKNPYEESS
	W.					RDVRSRRPADTEDGMSSCPQPWFVVIKEHQD
			İ			LKNGGQPVAGEDGQAADGSMQPTSWRQEPQ
						LCGMGTEQGCWIPVSSDKGSCPQVMERSFH
			1	l		MPSYGTQTLEGGVEKPHSLLSANPLWQQRAL
						DPPHQMELTQ
967	2317	A	8210	3	601	SSAMGSRSSHAAVIPDGDSIRRETGFSQASLL
			]	ļ		RLHHRFRALDRNKKGYLSRMDLQQIGALAV
						NPLGDRIIESFFPDGSQRVDFPGFVRVLAHFRP
						VEDEDTETQDPKKPEPLNSRRNKLHYAFQLY
1			1	ı	I	DLDRDGKISRHEMLQVLRLMVGVQVTEEQL
1			i			
						ENIADRTVQEADEDGDGAVSFVEFTKSLEKM DVEHKMSIRILK

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
968	2318	A	8211	2	409	ISSCPHTAYEGSMSTLSNFTQTLEDVFRRIFIT YMDNWRQNTTAEQEALQAKVDAENFYYVIL YLMVMIGMFSFIIVAILVSTVKSKRREHSNDP YHQYIVEDWQEKYKSQILNLEESKATIHENIG AAGFKMSP
969	2319	A	8215	1	1938	GMPRSRGGRAAPGPPPPPPPPPGQAPRWSRWR VPGRLLLLLPALCCLPGAARAAAAAAAGAGN RAAVAVAVARADEAEAPFAGQNWLKSYGY LLPYDSRASALHSAKALQSAVSTMQQFYGIP VTGVLDQTTIEWMKKPRCGVPDIIPHLSRRRR NKRYALTGQKWRQKHITYSIHNYTPKVGELD TRKAIRQAFDVWQKVTPLTFEEVPYHEIKSDR KEADIMIFFASGFHGDSSPFDGEGGFLAHAYF PGPGIGGDTHFDSDEPWTLGNANHDGNDLFL VAVHELGHALGLEHSSDPSAIMAPFYQYMET HNFKLPQDDLQGIQKIYGPPAEPLEPTRPLPTL PVRRIHSPSERKHERQPRPPRPPLGDRPSTPGT KPNICDGNFNTVALFRGEMFVFKDRWFWRL RNNRVQEGYPMQIEQFWKGLPARIDAAYER ADGRFVFFKGDKYWVFKEVTVEPGYPHSLG ELGSCLPREGIDTALRWEPVGKTYFFKGERY WRYSEERRATDPGYPKPITVWKGIPQAPQGA FISKEGYYTYFYKGRDYWKFDNQKLSVEPGY PRNILRDWMGCNQKEVERRKERRLPQDDVDI MVTINDVPGSVNAVAVVIPCILSLCILVLVYTI FQFKNKTGPQPVTYYKRPVQEWV
970	2320	A	8216	1235	2223	SRLSLQFYVSFRRTGLFTCKLIVEIFFRNYMN DSLRTNVFVRFQPETIACACIYLAARALQIPLP TRPHWFLLFGTTEEEIQEICIETLRLYTRKKPN YELLEKEVEKRKVALQEAKLKAKGLNPDGTP ALSTLGGFSPASKPSSPREVKAEEKSPISINVK TVKKEPEDRQQASKSPYNGVRKDSKRSRNSR SASRSRSRTRSRSRSHTPRRHYNNRRSRSGTY SSRSRSRSHSESPRRHHNHGSPHLKAKHTR DDLKSSNRHGHKRKKSRSRSQSKSRDHSDAA KKHRHERGHHRDRRERSRSFERSHKSKHHGG SRSGHGRHRR
971	2321	A	8217	3	3274	DCRLQAAMPTNFTVVPVEAHADGGGDETAE RTEAPGTPEGPEPERPSPGDGNPRENSPFLNN VEVEQESFFEGKNMALFEEEMDSNPMVSSLL NKLANYTNLSQGVVEHEEDEESRREAKAPR MGTFIGVYLPCLQNILGVILFLRLTWIVGVAG VLESFLIVAMCCTCTMLTAISMSAIATNGVVP AGGSYYMISRSLGPEFGGAVGLCFYLGTTFA GAMYILGTIEIFLTYISPGAAIFQAEAAGGEAA AMLHNMRVYGTCTLVLMALVVFVGVKYVN KLALVFLACVVLSILAIYAGVIKSAFDPPDIPV CLLGNRTLSRRSFDACVKAYGIHNNSATSAL WGLFCNGSQPSAACDEYFIQNNVTEIQGIPGA ASGVFLENLWSTYAHAGAFVEKKGVPSVPV AEESRASTLPYVLTDIAASFTLLVGIYFPSVTG IMAGSNRSGDLKDAQKSIPTGTILAIVTTSFIY LSCIVLFGACIEGVVLRDKFGEALQGNLVIGM LAWPSPWVIVIGSFFSTCGAGLQTLTGAPRLL QAIARDGIVPFLQVFGHGKANGEPTWALLLT VLICETGILIASLDSVAPILSMFFLMCYLFVNL ACAVQTLLRTPNWRPRFKFYHWTLSFLGMSL CLALMFICSWYYALSAMLIAGCIYKYIEYRG AEKEWGDGIRGLSLNAARYALLRVEHGPPHT KNWRPQVLVMLNLDAEQAMKHPRLLSFTSQ

ana	000	1.52	Lars	1 25	T	
SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence	ļ	09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence			914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
		İ	ĺ	amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
		ļ		residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
				peptide		/=possible nucleotide deletion, \=possible
		ļ	<b></b>	sequence		nucleotide insertion
						LKAGKGLTIVGSVLEGTYLDKHMEAQRAEE
						NIRSLMSTEKTKGFCQLVVSSSLRDGMSHLIQ
		ł	ł			SAGLGGLKHNTVLMAWPASWKQEDNPFSW
						KNFVDTVRDTTAAHQALLVAKNVDSFPQNQ
		l				ERFGGGHIDVWWIVHDGGMLMLLPFLLRQH
		}	1		1	KVWRKCRMRIFTVAQVDDNSIQMKKDLQMF
						LYHLRISAEVEVVEMVENDISAFTYERTLMM
			1			EQRSQMLKQMQLSKNEQEREAQLIHDRNTAS
		Ì	1			HTAAAARTQAPPTPDKVQMTWTREKLIAEK
						YRSRDTSLSGFKDLFSMKPDQSNVRRMIITAV
		1		1	4,0	KLNGVVLNKSQDAQLVLLNMPGPPKNRQGD
972	2322	<u> </u>	8224	701	246	ENYMEFLEVLTEGLNRVLLVRGGGREVITIYS
912	2322	A	8224	/01	240	TSRRVTMKFNPFVTSDRSKNRKRHFNAPSHV
	1	{			1	RRKIMSSPLSKELRQKYNVRSMPIRKDDEVQ
			1			VVRGHYKGQQIGKVVQVYRKKYVIYIERVQ
						REKANGTTVHVGIHPSKVVITRLKLDKDRKKI
973	2323	A	8237	873	4610	LERKAKSRQVGKEKGKYKEELIEKMQE
913	2323	A	8237	8/3	4010	GCPHAGGKGRVPTGGLTGGRTWSPSAAPRSC
ĺ		ĺ				PRPGPTPAPGAMDKLPPSMRKRLYSLPQQVG
			1			AKAWIMDEEEDAEEEGAGGRQDPSRRSIRLR
			i			PLPSPSPSAAAGGTESRSSALGAADSEGPARG
		ĺ	i		ĺ	AGKSSTNGDCRRFRGSLASLGSRGGGSGGTG SGSSHGHLHDSAEERRLIAEGDASPGEDRTPP
					1	GLAAEPERPGASAQPAASPPPPQQPPQPASAS
		<u> </u>				CEQPSVDTAIKVEGGAAAGDQILPEAEVRLG
		ł	ì		l	QAGFMQRQFGAMLQPGVNKFSLRMFGSQKA
						VEREQERVKSAGFWIIHPYSDFRFYWDLTML
			1		, , , , , , , , , , , , , , , , , , ,	LLMVGNLIIIPVGITFFKDENTTPWIVFNVVSD
l		ł	l			TFFLIDLVLNFRTGIVVEDNTEILDPQRIKMK
						YLKSWFMVDFISSIPVDYIFLIVETRIDSEVYK
ĺ	!	ĺ	1			TARALRIVRFTKILSLLRLLRLSRLIRYIHQWE
ļ						EIFHMTYDLASAVVRIVNLIGMMLLLCHWDG
						CLQFLVPMLQDFPDDCWVSINNMVNNSWGK
Í	ı	ĺ	1	·		QYSYALFKAMSHMLCIGYGRQAPVGMSDV
						WLTMLSMIVGATCYAMFIGHATALIOSLDSS
ļ						RRQYQEKYKQVEQYMSFHKLPPDTRQRIHD
		ĺ	ì			YYEHRYQGKMFDEESILGELSEPLREEIINFNC
			1 .			RKLVASMPLFANADPNFVTSMLTKLRFEVFQ
						PGDYIIREGTIGKKMYFIQHGVVSVLTKGNKE
ł			ì			TKLADGSYFGEICLLTRGRRTASVRADTYCR
						LYSLSVDNFNEVLEEYPMMRRAFETVALDRL
						DRIGKKNSILLHKVQHDLNSGVFNYQENEIIQ
						QIVQHDREMAHCAHRVQAAASATPTPTPVIW
			1			TPLIQAPLQAAAATTSVAIALTHHPRLPAAIFR
				ĺ		PPPGSGLGNLGAGOTPRHLKRLOSLIPSALGS
						ASPASSPSQVDTPSSSSFHIQQLAGFSAPAGLS
						PLLPSSSSSPPPGACGSPSAPTPSAGVAATTIA
1						GFGHFHKALGGSLSSSDSPLLTPLQPGARSPQ
						AAQPSPAPPGARGGLGLPEHFLPPPPSSRSPSS
						SPGQLGQPPGELSLGLATGPLSTPETPPRQPEP
1		-				PSLVAGASGGASPVGFTPRGGLSPPGHSPGPP
						RTFPSAPPRASGSHGSLLLPPASSPPPPQVPQR
í				[		RGTPPLTPGRLTQDLKLISASQPALPQDGAQT
I						LRRASPHSSGESMAAFPLFPRAGGGSGSGSSS
				I I		GGLGPPGRPYGAIPGQHVTLPRKTSSGSLPPP
			1			OCCUPATOR LOWINGOUS ITERN 19909PAR
						LSLFGARATSSGGPPLTAGPQREPGARPEPVR SKLPSNL
974	2324	A	8247	279	468	LSLFGARATSSGGPPLTAGPQREPGARPEPVR SKLPSNL
974	2324	A	8247	279	468	LSLFGARATSSGGPPLTAGPQREPGARPEPVR

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
975	2325	A	8249	62	1571	LVALKNWKPKGTNIPAPQSPVFGEAVSGVYM MTKVLGMAPVLGPRPPQEQVGPLMVKVEEK EEKGKYLPSLEMFRQRFRQFGYHDTPGPREA LSQLRVLCCEWLRPEIHTKEQILELLVLEQFLT ILPQELQAWVQEHCPESAEEAVTLLEDLEREL DEPGHQVSTPPNEQKPVWEKISSSGTAKESPS SMQPQPLETSHKYESWGPLYIQESGEEQEFAQ DPRKVRDCRLSTQHEESADEQKGSEAEGLKG DIISVIIANKPEASLERQCVNLENEKGTKPPLQ EAGSKKGRESVPTKPTPGERRYICAECGKAFSNSSNLTKHRRTHTGEKPYVCTKCGKAFSHSS NLTLHYRTHLVDRPYDCKCGKAFSGKGSLIRHYR IHTGEKPYQCNECGKSFSQHAGLSSHQRLHT GEKPYKCKECGKAFNHSSNFNKHHRIHTGEK PYWCHHCGKTFCSKSNLSKHQRVHTGEGEA P
976	2326	A	8257	298	7086	GNMACWPQLRLLLWKNLTFRRRQTCQLLLE VAWPLFIFLILISVRLSYPPYEQHECHFPNKAM PSAGTLPWVQGIICNANNPCFRYPTPGEAPGV VGNFNKSIVARLFSDARRLLLYSQKDTSMKD MRKVLRTLQQIKKSSSNLKLQDFLVDNETFS GFLYHNLSLPKSTVDKMLRADVILHKVFLQG YQLHLTSLCNGSKSEEMIQLGDQEVSELCGLP REKLAAAERVLRSNMDILKPILRITLNSTSPFPS KELAEATKTLLHSLGTLAQELFSMRSWSDMR QEVMFLTNVNSSSSSTQIYQAVSRIVCGHPEG GGLKIKSLNWYEDNNYKALFGGNGTEEDAE TFYDNSTTPYCNDLMKNLESSPLSRIIWKALK PLLVGKILYTPDTPATRQVMAEVNKTFQELA VFHDLEGMWEELSPKIWTFMENSQEMDLVR MLLDSRDNDHFWEQQLDGLDWTAQDIVAFL AKHPEDVQSSNGSVYTWREAFNETNQAIRTIS RFMECVNLNKLEPIATEVWLINKSMELLDER KFWAGIVFTGITPGSIELPHHVKYKIRMGIDN VERTNKIKDGYWDPGPRADPFEDMRYVWGG FAYLQDVVEQAIIRVLTGTEKKTGVYMQQMP YPCYVDDIFLRVMSRSMPLFMTLAWIYSVAV IIKGIVYEKEARLKETMRIMGLDNSILWFSWFI SSLIPLLVSAGILVVILKLGNLLPYSDPSVVFV FLSVFAVVTILQCFLISTLFSRANLAAACGGII YFTLYLPYVLCVAWQDYVGFTLKIFASLLSP VAFGFGCEYFALFEEQGIGVQWDNLFESPVE EDGFNLTTSVSMMLFDTFLYGVMTWYIEAVF PGQYGIPRPWYFPCTKSYWFGEESDEKSHPGS NQKRISEICMEEEPTHLKLGVSIQNLVKVYRD GMKVAVDGLALNFYEGQITSFLGHNGAGKT TTMSILTGLFPPTSGTA YILGKDRISEMSTIRQ NLGVCPQHNVLFDMLTVEEHIWFYARLKGLS EKHVKAEMEQMALDVGLPSSKLKSKTSQLS GGMQRKLSVALAFVGGSKVVILDEPTAGVDP YSRRGIWELLLKYRQGRTIILSTHHMDEADVL GDRIAIISHGKLCCVGSSLFLKNQLGTGYYLT LVKKDVESSLSSCRNSSSTVSYLKKEDSVSQS SSDAGLGSDHESDTLTIDVSAISNLIRKHVSEA RLVEDIGHELTYVLPYEAAKEGAFVELFHEID DRLSDLGISSYGISETTLEEIFLKVAEESGVDA ETSDGTLPARRNRAFGDKQSCLRPFTEDDA ADPNDSDIDPESRETDLLSGMDGKGSYQVKG WKLTQQGFVALLWKRLLIARRSRKGFFAQIV

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Scrine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
						LPAVFVCIALVFSLIVPPFGKYPSLELQPWMY NEQYTFVSNDAPEDTGTLELLNALTKDPGFG TRCMEGNPIPDTPCQAGEEEWTTAPVPQTIM DLFQNGNWTMQNPSPACQCSSDKIKKMLPV CPPGAGGLPPPQRKQNTADILQDLTGRNISDY LVKTYVQIIAKSLKNKIWVNEFFYGGFSLGVS NTQALPPSQEVNDATKQMKKHLKLAKDSSA DRFLNSLGRFMTGLDTRNNVKVWFNNKGW HAISSFLNVINNAILRANLQKGENPSHYGITAF NHPLNLTKQQLSEVAPMTTSVDVLVSICVIFA MSFVPASFVVFLIQERVSKAKHLQFISGVKPVI YWLSNFVWDMCNYVVPATLVIIIFICFQQKSY VSSTNLPVLALLLLLYGWSITPLMYPASFVFK IPSTAYVVLTSVNLFIGINGSVATFVLELFTDN KLNNINDILKSVFLIFPHFCLGRGLIDMVKNQ AMADALERFGENRFVSPLSWDLVGRNLFAM AVEGVVFFLITVLIQYRFIRPRPVNAKLSPLN DEDEDVRRERQRILDGGGQNDILEIKELTKIY RRKRKPAVDRICVGIPPGECFGLLGVNGAGK SSTFKMLTGDTTVTRGDAFLNRNSILSNIHEV HQNMGYCPQFDAITELLTGREHVEFFALLRG VPEKEVGKVGEWAIRKLGLVKYGEKYAGNY SGGNKKKLSTAMALIGGPPVVFLDEPTTGMD PKARRFLWNCALSVVKEGRSVVLTSHSMEEC EALCTRMAIMVNGRFRCLGSVQHLKNRFGD GYTIVVRIAGSNPDLKPVQDFFGLAFPGSVPK EKHRNMLQYQLPSSLSSLARIFSILSQSKKRLH IEDYSVSQTTLDQVFVNFAKDQSDDDHLKDL SLHKNQTVVDVAVLTSFLQDEKVKESYV
977	2327	A	8260	3	1567	IPGSTISFSLCFIFPPCVPTMVRKPVVSTISKGG YLQGNVNGRLPSLGNKEPPGQEKVQLKRKV TLLRGVSIIIGTIIGAGIFISPKGVLQNTGSVGM SLTIWTVCGVLSLFGALSYAELGTTIKKSGGH YTYILEVFGPLPAFVRVWVELLIIRPAATAVIS LAFGRYILEPFFIQCEIPELAIKLITAVGITVVM VLNSMSVSWSARIQIFLTFCKLTAILIIIVPGV MQLIKGQTQNFKDAFSGRDSSITRLPLAFYYG MYAYAGWFYLNFVTEEVENPEKTIPLAICISM AIVTIGYVLTNVAYFTTINAEELLLSNAVAVT FSERLLGNFSLAVPIFVALSCFGSMNGGVFAV SRLFYVASREGHLPEILSMIHVRKHTPLPAVIV LHPLTMIMLFSGDLDSLLNFLSFARWLFIGLA VAGLIYLRYKCPDMHRPFKVPLFIPALFSFTC LFMVALSLYSDPFSTGIGFVITLTGVPAYYLFII WDKKPRWFRIMSEKITRTLQIILEVVPEEDKL
978	2328	A	8261	2	2165	RGGSLRCVLGKLLGQLLCFQSERCVRFPEGLL RHRGCGLLSSRLSAGKPPLRTSFFGSWGVLPP LADAASMSGVRAVRISIESACEKQVHEVGLD GTETYLPPLSMSQNLARLAQRIDFSQGSGSEE EEAAGTEGDAQEWPGAGSSADQDDEEGVVK FQPSLWPWDSVRNNLRSALTEMCVLYDVLSI VRDKKFMTLDPVSQDALPPKQNPQTLQLISK KKSLAGAAQILLKGAERLTKSVTENQENKLQ RDFNSELLRLRQHWKLRKVGDKILGDLSYRS AGSLFPHHGTFEVIKNTDLDLDKKIPEDYCPL DVQIPSDLEGSAYIKVSIQKQAPDIGDLGTVN LFKRPLPKSKPGSPHWQTKLEAAQNVLLCKEI FAQLSREAVQIKSQVPHIVVKNQIISQPFPSLQ LSISLCHSSNDKKSQKFATEKQCPEDHLYVLE HNLHLLIREFHKQTLSSIMMPHPASAPFGHKR

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979	2329	A	8289	2	1053	WNKMEGRNFVYKMELLMSALSPCLL FVWNPRGGRKRRRQAAVTQAATRASGTPSP RDGTMTQGKLSVANKAPGTEGQQQVHGEKK EAPAVPSAPPSYEEATSGEGMKAGAFPAPTA VPLHPSWAYVDPSSSSSYDNGFPTGDHELFTT FSWDDQKVRRVFVRKVYTILLIQLLVTLAVV ALFTFCDPVKDYVQANPGWYWASYAVFFAT YLTLACCSGPRRHFPWNLILLTVFTLSMAYLT GMLSSYYNTTSVLLCLGITALVCLSVTVFSFQ TKFDFTSCQGVLFVLLMTLFFSGLILAILLPFQ YVPWLHAVYAALGAGVFTLFLALDTQLLMG NRRHSLSPEEYIFGALNIYLDIIYIFTFFLQLFG TNRE
980	2330	A	8305	59	857	ASQLPDYSISPPSLPPRISFHPSPTLARVAMAEP SEATQSHSISSSSFGAEPSAPGGGSPGACPAL GTKSCSSSCAVHDLIFWRDVKKTGFVFGTTLI MLSLAAFSVISVVSYLILALLSVTISFRIYKSV IQAVQKSEEGHPFKAYLDVDITLSSEAFHNY MNAAMVHINRALKLIIRLFLVEDLVDSLKLA VFMWLMTYVGAVFNGITLLILAELLIFSVPIV YEKYKTQIDHYVGIARDQTKSIVEKIQAKLPG IAKKKAE
981	2331	A	8308	186	1337	TRMSRHEGVSCDACLKGNFRGRRYKCLICYD YDLCASCYESGATTTRHTTDHPMQCILTRVD FDLYYGGEAFSVEQPQSFICPYCGKMGYTET SLQEHVTSEHAETSTEVICPICAALPGGDPNH VTDDFAAHLTLEHRAPRDLDESSGVRHVRR MFHPGRGLGGPRARRSNMHFTSSSTGGLSSS QSSYSPSNREAMDPIAELLSQLSGVRRSAGGQ LNSSGPSASQLQQLQMQLQLERQHAQAARQ QLETARNATRRTNTSSVITTITQSTATTNIAN TESSQQTLQNSQFLLTRLNDPKMSETERQSM ESERADRSLFVQELLLSTLVREESSSDEDDR GEMADFGAMGCVDIMPLDVALENLNLKESN KGNEPPPPPL
982	2332	A	8315	1	1004	GSTHASADAWAQWFCTEALVMGAPVWYLV AAALLVGFILFLTRSRGRAASAGQEPLHNEEL AGAGRVAQPGPLEPEEPRAGGRPRRRDLGS RLQAQRRAQRVAWAEADENEEEAVILAQEE EGVEKPAETHLSGKIGAKKLRKLEEKQARKA QREAEEAEREERKRLESQREAEWKKEEERLR LEEEQKEEEERKAREEQAQREHEEYLKLKEA FVVEEEGVGETMTEEQSQSFLTEFINYIKQSK VVLLEDLASQVGLRTQDTINRIQDLLAEGTIT GVIDDRGKFIYITPEELAAVANFIRQRGRVSIA ELAQASNSLIAWGRESPAQAPA
983	2333	A	8320	244	1420	RRRWRARGGLVPTLAWAEATGAYVPGRDKP DLPTWKRNFRSALNRKEGLRLAEDRSKDPHD PHKIYEFVNSGVGDFSQPDTSPDTNGGGSTSD TQEDILDELLGNMVLAPLPDPGPPSLAVAPEP CPQPLRSPSLDNPTPFPNLGPSENPLKRLLVPG EEWEFEVTAFYRGRQVFQQTISCPEGLRLVGS

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984	2334	A	8321	1	1243	ANMAPVEHVVADAGAFLRHAALQDIGKNIY TIREVVTEIRDKATRRLAVLPYELRFKEPLPE YVRLVTEFSKKTGDYPSLSATDIQVLALTYQL EAEFVGVSHLKQEPQKVKVSSSIQHPETPLHIS GFHLPYKPKPPQETEKGHSACEPENLEFSSFM FWRNPLPNIDHELQELLIDRGEDVPSEEEEEE NGFEDRKDDSDDDGGGWITPSNIKQIQQELE QCDVPEDVRVGCLTTDFAMQNVLLQMGLHV LAVNGMLIREARSYILRCHGCFKTTSDMSRV FCSHCGNKTLKKVSVTVSDDGTLHMHFSRNP KVLNPRGLRYSLPTPKGGKYAINPHLTEDQRF PQLRLSQKARQKTNVFAPDYIAGVSPFVENDI SSRSATLQVRDSTLGAGRRRLNPNASRKKFV KKR
985	2335	A	8322	352	529	RRNNIRQFIMKVCISGQARWLTPVVPVLWET EAGRSLELKSLRPAWATWGNPISTKINK
986	2336	A	8325	89	1172	KMNPTDIADTTLDESIYSNYYLYESIPKPCTKE GIKAFGELFLPPLYSLVFVFGLLGNSVVVLVL FKYKRLRSMTDVYLLNLAISDLLFVFSLPFWG YYAADQWVFGLGLCKMISWMYLVGFYSGIF FVMLMSIDRYLAIVHAVFSLRARTLTYGVITS LATWSVAVFASLPGFLFSTCYTERNHTYCKT KYSLNSTTWKVLSSLEINILGLVIPLGIMLFCY SMIIRTLQHCKNEKKNKAVKMIFAVVVLFLG FWTPYNIVLFLETLVELEVLQDCTFERYLDYA IQATETLAFVHCCLNPIIYFFLGEKFRKYILQL FKTCRGLFVLCQYCGLLQIYSADTPSSSYTQS TMDHDLHDAL
987	2337	A	8326	3	470	SLSAMRFLAATFLLLALSTAAQAEPVQFKDC GSVDGVIKEVNVSPCPTQPCQLSKGQSYSVN VTFTSNIQSKSSKAVVHGILMGVPVPFPIPEPD GCKSGINCPIQKDKTYSYLNKLPVKSEYPSIK LVVEWQLQDDKNQSLFCWEIPVQIVSHL
988	2338	A	8335	1205	323	VIKMALAARLLPQFLHSRSLPCGAVRLRTPA VAEVRLPSATLCYFCRCRLGLGAALFPRSAR ALAASALPAQGSRWPVLSSPGLPAAFASFPAC PQRSYSTEEKPQQHQKTKMIVLGFSNPINWV RTRIKAFLIWAYFDKEFSITEFSEGAKQAFAH VSKLLSQCKFDLLEELVAKEVLHALKEKVTS LPDNHKNALAANIDEIVFTSTGDISIYYDEKG RKFVNILMCFWYLTSANIPSETLRGASVFQVK LGNQNVETKQLLSASYEFQREFTQGVKPDWT IARIEHSKLLE
989	2339	A	8349	67	185	MSGFIHQLLIQNLFCVYHTRLKTSQGLCLLSL KSLHPMS
990	2340	A	8361	210	1115	ASPFLRPQGHDSGEREPFSQTPGLMQPFSIPVQ ITLQGSRRRQGRTAFPASGKKRETDYSDGDPL DVHKRLPSSTGEDRAVMLGFAMMGFSVLMF FLLGTTILKPFMLSIQREESTCTAIHTDIMDDW LDCAFTCGVHCHGQGKYPCLQVFVNLSHPG QKALLHYNEEAVQINPKCFYTPKCHQDRNDL LNSALDIKEFFDHKNGTPFSCFYSPASQSEDVI

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991	2341	A	8369	9	921	SSVVEFSALSVSMACLSPSQLQKFQQDGFLVL EGFLSAEECVAMQQRIGEIVAEMDVPLHCRT EFSTQEEEQLRAQGSTDYFLSSGDKIRFFFEK GVFDEKGNFLVPPEKSINKIGHALHAHDPVFK SITHSFKVQTLARSLGLQMPVVVQSMYIFKQP HFGGEVSPHQDASFLYTEPLGRVLGVWIAVE DATLENGCLWFIPGSIITSGVSRRMVRAPVGS APGTSFLGSEPARDNSLFVPTPVQRGALVLIH GEVVHKSKQNLSDRSRQAYTFHLMEASGTT WSPENWLQPTAELPFPQLYT
992	2342	A	8370	906	4	MALSGNCSRYPPREQGSAVPNSFPEVVELNV GGQVYFTRHSTLISIPHSLLWKMFSPKRDTAN DLAKDSKGRFFIDRDGFLFRYILDYLRDRQVV LPDHFPEKGRLKREAEYFQLPDLVKLLTPDEI KQSPDEFCHSDFEDASQGSDTRICPPSSLLPAD RKWGFITVGYRGSCTLGREGQADAKFRRVPR ILVCGRISLAKEVFGETLNESRDPDRAPERYTS RFYLKFKHLMGAPASNFILGFWGLGQNQDK HPVNIYLQQRSVIRPDLTSKKAGDLKGKGDA QEVSRRRWLGDPEHL
993	2343	A	8379	1	2794	MRMQRHKNDTMDFGDSGKRIGGGVLCLLHQ SNTSFIKLNNNGFEDIVIVIDPSVPEDEKIIEQIE DMVTTASTYLFEATEKRFFFKNVSILIPENWK ENPQYKRPKHENHKHADVIVAPPTLPGRDEP YTKQFTECGEKGEYIHFTPDLLLGKKQNEYG PPGKLFVHEWAHLRWGVFDEYNEDQPFYRA KSKKIEATRCSAGISGRNRVYKCQGGSCLSRA CRIDSTTKLYGKDCQFFPDKVQTEKASIMFM QSIDSVVEFCNEKTHNQEAPSLQNIKCNFRST WEVISNSEDFKNTIPMVTPPPPPVFSLLKIRQRI VCLVLDKSGSMGGKDRLNRMNQAAKHFLLQ TVENGSWVGMVHFDSTATIVNKLIQIKSSDER NTLMAGLPTYPLGGTSICSGIKYAFQVIGELH SQLDGSEVLLLTDGEDNTASSCIDEVKQSGAI VHFIALGRAADEAVIEMSKITGGSHFYVSDEA QNNGLIDAFGALTSGNTDLSQKSLQLESKGLT LNSNAWMNDTVIIDSTVGKDTFFLITWNSLPP SISLWDPSGTIMENFTVDATSKMAYLSIPGTA KVGTWAYNLQAKANPETLTITVTSRAANSSV PPITVNAKMNKDVNSFPSPMIVYAEILQGYVP VLGANVTAFIESQNGHTEVLELLDNGAGADS FKNDGVYSRYFTAYTENGRYSLKVRAHGGA NTARLKLRPPLNRAAYIPGWVVNGEIEANPP RPEIDEDTQTTLEDFSRTASGGAFVVSQVPSL PLPDQYPPSQITDLDATVHEDKIILTWTAPGD NFDVGKVQRYIIRISASILDLRDSFDDALQVN TTDLSPKEANSKESFAFKPENISEENATHIFIAI KSIDKSNLTSKVSNIAQVTLFIPQANPDDIDPT PTPTPTPTPDKSHNSGVNISTLVLSVIGSVVIV
994	2344	A	8385	231	644	INSSPRTGRDHQELNLHTERDSRSQRAVLKIP RQNPGIFYWIFLPSRSHSASHGSRQRQVSCQG TQDEILKMRNTFAELKNSLEALSSRMDQAEE RIGTQAGVQWRDHGSLQPQPPEFKQCFHLSL
995	2345	A	8390	194	3421	PSSWDYRACLS AWRKSSVVPPRGTRRGEKSDQDKSGQKNKR

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SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide	ļ	in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence			914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
				amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
				peptide		/=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion
						DFLSMKQSPALAPEERCRRAGSPKPVLRADD
						NNMGNGCSQKLATANLLRFLLLVLIPCICALV
						LLLEILLSYVGTLQKVYFKSNGSEPLVTDGEI
		1				OGSDVILTNTIYNQSTVVSTAHPDQHVPAWT
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	i	l	ł	ĺ		ATLTPLLSVVRNMEMEKFLKFFTYLHRLSCY
						QHIMLFGCTLAFPECIIDGDDSHGLLPCRSFCE
						AAKEGCESVLGMVNYSWPDFLRCSQFRNQT
						ESSNVSRICFSPQQENGKQLLCGRGENFLCAS
	1					GICIPGKLQCNGYNDCDDWSDEAHCNCSENL
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						DCNPTTEHRCGDGRCIAMEWVCDGDHDCVD
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						KSDEVNCSCHSQGLVECRNGQCIPSTFQCDG
						DEDCKDGSDEENCSVIQTSCQEGDQRCLYNP
1						CLDSCGGSSLCDPNNSLNNCSQCEPITLELCM
	1	1	1	1		NLPYNSTSYPNYFGHRTQKEASISWESSLFPA
						LVQTNCYKYLMFFSCTILVPKCDVNTGEHIPP
1	1					CRALCEHSKERCESVLGIVGLQWPEDTDCSQ
						FPEENSDNQTCLMPDEYVEECSPSHFKCRSGQ
						CVLASRRCDGQADCDDDSDEENCGCKERDL
						WECPSNKQCLKHTVICDGFPDCPDYMDEKN
	J			j		CSFCQDDELECANHACVSRDLWCDGEADCS
		1				DSSDEWDCVTLSINVNSSSFLMVHRAATEHH
						VCADGWQEILSQLACKQMGLGEPSVTKLIQE
						QEKEPRWLTLHSNWESLNGTTLHELLVNGQS
						CESRSKISLLCTKQDCGRRPAARMNKRILGGR
						TSRPGRWPWQCSLQSEPSGHICGCVLIAKKW
						VLTVAHCFEGRENAAVWKVVLGINNLDHPS
Ĭ	1	İ	(	1		VFMQTRFVKTIILHPRYSRAVVDYDISIVELSE
						DISETGYVRPVCLPNPEQWLEPDTYCYITGW
						GHMGNKMPFKLQEGEVRIISLEHCQSYFDMK
1						TITTRMICAGYESGTVDSCMGDSGGPLVCEK
1						PGGRWTLFGLTSWGSVCFSKVLGPGVYSNVS
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996	2346	A	8392	199	3085	KVILSSEMSKTNKSKSGSRSSRSRSASRSRSRS
	2340	Α	0372	1//	3003	FSKSRSRSRSLSRSRKRRLSSRSRSRSYSPAHN
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						KDSRPSQAAGDNQGDEVKEQTFSGGTSQDTK
	1					ASESSKPWPDATYGTGSASRASAVSELSPRER
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1	[					KGSFSDTGLGDGKMKSDSFAPKTDSEKPFRG
1	-					SQSPKRYKLRDDFEKKMADFHKEEMDDQDK
						DKAKGRKESEFDDEPKFMSKVIGANKNQEEE
1	1					KSGKWEGLVYAPPGKEKQRKTEELEEESFPE
I	J	J	J	]	]	RSKKEDRGKRSEGGHRGFVPEKNFRVTAYK
1						AVQEKSSSPPPRKTSESRDKLGAKGDFPTGKS
1						SFSITREAQVNVRMDSFDEDLARPSGLLAQER
[		1				KLCRDLVHSNKKEQEFRSIFQHIQSAQSQRSP
						SELFAQHIVTIVHHVKEHHFGSSGMTLHERFT
						KYLKRGTEQEAAKNKKSPEIHRRIDISPSTFRK
1						HGLAHDEMKSPREPGYKAEGKYKDDPVDLR
1				4		, The transfer of the contract of the co
	ľ	[	( .	1		LDIERRKKHKERDLKRGKSRESVDSRDSSHSR
						LDIERRKKHKERDLKRGKSRESVDSRDSSHSR ERSAEKTEKTHKGSKKOKKHRRARDRSRSSS
		ļ				LDIERRKKHKERDLKRGKSRESVDSRDSSHSR ERSAEKTEKTHKGSKKQKKHRRARDRSRSSS SSSQSSHSYKAEEYTEETEEREESTTGFDKSRL

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						GTKDFVGPSERGGGRARGTFQFRARGRGWG RGNYSGNNNNNSNNDFQKRNREEEWDPEYT PKSKKYYLHDDREGEGSDKWVSRGRGRGAF PRGRGRFMFRKSSTSPKWAHDKFSGEEGEIE DDESGTENREEKDNIQPTTE
997	2347	A	8398	202	552	CPALGGRQDLQGTRLLWAHDSGVGGQKAKS KQENLESLEATGREEEGGQGPPVTTKGVLLA LLMAGLALQPGTALLCYSCKAQVSNEDCLQ VENCTQLGEQCWTARIREWGDDSRQA
998	2348	A	8400	697	301	NPPSACTPGSCDSCSGRGRDLAFDSVWSTNN MSDPRRPNKVLRYKPPPSECNPALDDPTPDY MNLLGMIFSMCGLMLKLKWCAWVAVYCSFI SFANSRSSEDTKQMMSSFMLSISAVVMSYLQ NPQPMTPPW
999	2349	A	8401	93	1126	ASASHITSGHLRCFPGSEGVGTMARCFSLVLL LTSIWTTRLLVQGSLRAEELSIQVSCRIMGITL VSKKANQQLNFTEAKEACRLLGLSLAGKDQ VETALKASFETCSYGWVGDGFVVISRISPNPK CGKNGVGVLIWKVPVSRQFAAYCYNSSDTW TNSCIPEIITTKDPIFNTQTATQTTEFIVSDSTYS VASPYSTIPAPTTTPPAPASTSIPRRKKLICVTE VFMETSTMSTETEPFVENKAAFKNEAAGFGG VPTALLVLALLFFGAAAGLGFCYVKRYVKAF PFTNKNQQKEMIETKVVKEEKANDSNPNEES KKTDKNPEESKSPSKTTMRCLEAEV
1000	2350	A	8406	2	777	KERCQFVVKPMLSTVGSFLQDLQNEDKGIKT AAIFTADGNMISASTLMDILLMNDFKLVINKI AYDVQCPKREKPSNEHTAEMEHMKSLVHRL FTILHLEESQKKREHHLLEKIDHLKEQLQPLE QVKAGIEAHSEAKTSGLLWAGLALLSIQGGA LAWLTWWVYSWDIMEPVTYFITFANSMVFF AYFIVTRQDYTYSAVKSRQFLQFFHKKSKQQ HFDVQQYNKLKEDLAKAKESLKQARHSLCL QMQVEELNEKN
1001	2351	A	8410	1400		VGFWERPLRSSRWFRRSLRRWEMLARAARG TGALLLRGSLLASGRAPRRASSGLPRNTVVLF VPQQEAWVVERMGRFHRILEPGLNILIPVLDR IRYVQSLKEIVINVPEQSAVTLDNVTLQIDGV LYLRIMDPYKASYGVEDPEYAVTQLAQTTM RSELGKLSLDKVFRERESLNASIVDAINQAAD CWGIRCLRYEIKDIHVPPRVKESMQMQVEAE RRKRATVLESEGTRESAINVAEGKKQAQILAS EAEKAEQINQAAGEASAVLAKAKAKAEAIRI LAAALTQHNGDAAASLTVAEQYVSAFSKLA KDSNTILLPSNPGDVTSMVAQAMGVYGALT KAPVPGTPDSLSSGSSRDVQGTDASLDEELDR VKMS
1002	2352	A	8421	134	941	NRENLLESRMMDPCSVGVQLRTTNECHKTY YTRHTGFKTLQELSSNDMLLLQLRTGMTLSG NNTICFHHVKIYIDRFEDLQKSCCDPFNIHKKL AKKNLHVIDLDDATFLSAKFGRQLVPGWKLC PKCTQIINGSVDVDTEDRQKRKPESDGRTAK ALRSLQFTNPGRQTEFAPETGKREKRRLTKN ATAGSDRQVIPAKSKVYDSQGLLIFSGMDLC DCLDEDCLGCFYACPACGSTKCGAECRCDRK WLYEQIEIEGGEIIHNKHAG
1003	2353	A	8427	3	1416	TEWGLSGSCPGCSPLEPGSRGRGAAAWRILR CRRLPEPSPFLTQPNLAQSQPPAPVPVTDPSVT MHPAVFLSLPDLRCSLLLLVTWVFTPVTTEIT

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						PESSFQKLAPSEYRYTLLRDRDEL
1004	2354.	A	8432	910	387	GLSRKLRAGFLPGFCRVSPCGSWVVETLVKM ACAAARSPADQDRFICIYPAYLNNKKTIAEGR RIPISKAVENPTATEIQDVCSAVGLNVFLEKN KMYSREWNRDVQYRGRVRVQLKQEDGSLC LVQFPSRKSVMLYAAEMIPKLKTRTQKTGGA DQSLQQGEGSKKGKGKKKK
1005	2355	A	8453	90	530	QSHETKMQSGTHWRVLGLCLLSVGVWGQD GNEEMGGITQTPYKVSISGTTVILTCPQYPGSE ILWQHNDKNIGGDEDDKNIGSDEDHLSLKEF SELEQSGYYVCYPRGSKPEDANFYLYLRARG NPGLQNRYHRLFREDHSKGHSQ
1006	2356	A	8458	3	307	AVQRIRHEMNIFRLTGDLSHLAAIVILLLKIW KTRSCAGISGKSQLLFALVFTTRYLDLFTSFIS LYNTSMKVWYAIHRNVFHLQCTGLWTLNLC QLCIFN
1007	2357	A	8459	43	553	GAGAGGDWAAMDKLKKVLSGQDTEDRSGL SEVVEASSLSWSTRIKGFIACFAIGILCSLLGT VLLWVPRKGLHLFAVFYTFGNIASIGSTIFLM GPVKQLKRMFEPTRLIATIMVLLCFALTLCSA FWWHNKGLALIFCILQSLALTWYSLSFIPFAR DAVKKCFAVCLA
1008	2358	Ā	8462	487	150	AQDIRSVHSLGQKSTFVKHFRTLSHLHGLPDP PPHWPPQERSPPSHPCMPSHRPQIPQLSNSGPS DPRWGCVGPSMPTSTCLPGAVEASTTKASLP KCPVDSSLPTPEACFL
1009	2359	A	8465	134	954	ETRVKTSLELLRTQLEPTGTVGNTIMTSQPVP NETIIVLPSNVINFSQAEKPEPTNQGQDSLKKH LHAEIKVIGTIQILCGMMVLSLGIILASASFSPN FTQVTSTLLNSAYPFIGPFFFIISGSLSIATEKRL TKLLVHSSLVGSILSALSALVGFIILSVKQATL NPASLQCELDKNNIPTRSYVSYFYHDSLYTTD CYTAKASLAGTLSLMLICTLLEFCLAVLTAVL RWKQAYSDFPGSVLFLPHSYIGNSGMSSKMT HDCGYEELLTS
1010	2360	A	8468	2	473	KYRYRRPYPVMRKICQVGPAGLAFILNISPVA HRVALCHLAGCQEQAAWYHTLQILFFLVSAY FFSCPVPEKYFPGSCDIVGHGHQIFHAFLSICT LSQLEAILLDYQGRQEIFLQRHGPLSVHMACL SFFFLAACSAATAALLRHKVKARLTKKDS
1011	2361	A	8478	5	409	TELSQLEKAHPPADMGRRKSKRKPPPKKKMT GTLETQFTCPFCNHEKSCDVKMDRARNTGVI SCTVCLEEFQTPITCILGNLGFFQRVGRGLESG PCSSGPLCALVQGQSRPEEQVPPSDFCGVRRC RAGFQCQ
1012	2362	A	8481	2810	1652	RTSTQKWQSVFNDSQEHLERFYCNPENDRM RMKYGGQEFWADLNAMNVYETTEFDQLRR LSTPPSSNVNSIYHTVWKFFCRDHFGWREYPE

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						VTSDLYDSCVDNFFEPQIFVIFNDDQSYPYFVI QYEEVSNTVSI
1013	2363	A	8488	2	517	IENCRTRLRQAWHEVCGNKMAAPIPQGFSCL SRFLGWWFRQPVLVTQSAAIVPVRTKKRFTP PIYQPKFKTEKEFMQHARKAGLVIPPEKSDRS IHLACTAGIFDAYVPPEGDARISSLSKEGLIER TERMKKTMASQVSIRRIKDYDANFKIKDFPE KAKDIFIEGSPLY
1014	2364	. A	8501	363	17	YIRTGYVYICIIYAQLMYTYYIRTAYVYICILY AQLMYTYVLYTHSLCIHMYSIRTAYVYICIIY AQIMYTYVFYTHRLCIHMYSIRTDYVYICILY AQLMYTYVFYTHSYMSDE
1015	2365	Α	8504	3	2190	NSSEHFSQAPQRLSFYSWYGSARLFRFRVPPD AVLLRWLLQVSRESGAACTDAEITVHFRSGA PPVINPLGTSFPDDTAVQPSFQVGVPLSTTPRS NASVNVSHPAPGDWFVAAHLPPSSQKIELKG LAPTCAYVFQPELLVTRVVEISIMEPDVPLPQ TLLSHPSYLKVFVPDYTRELLLELRDCVSNGS LGCPVRLTVGPVTLPSNFQKVLTCTGAPWPC RLLLPSPPWDRWLQVTAESLVGPLGTVAFSA VAALTACRPRSVTIQPLLQSSQNQSFNASSGL LSPSPDHQDLGRSGRVDRSPFCLTNYPVTRED MDVVSVHFQPLDRVSVRVCSDTPSVMRLRL NTGMDSGGSLTISLRANKTEMRNETVVVACV NAASPFLGFNTSLNCTTAFFQGYPLSLSAWSR RANLIIPYPETDNWYLSLQLMCPENAEDCEQ AVVHVETTLYLVPCLNDCGPYGQCLLLRRHS YLYASCSCKAGWRGWSCTDNSTAQTVAQQR AATLLLTLSNLMFLAPIAVSVRFFLVEASVY AYTMFFSTFYHACDQPGEAVLCILSYDTLQY CDFLGSGAAIWVTILCMARLKTVLKYVLFLL GTLVIAMSLQLDRRGMWNMLGPCLFAFVIM ASMWAYRCGHRRQCYPTSWQRWAFYLLPG VSMASVGIAIYTSMMTSDNYYYTHSIWHILL AGSAALLLPPPDQPAEPWACSQKFPCHYQIC KNDREELYAVT
1016	2366	A	8511	1	453	KWYPSGPVRIPGRFYYKLPAGHRRCRMAPAK KGGEKKKGRSAINEVVTREYTINIHKRIHGVG FKKRAPRALKEIRKFAMKEMGTPDVRIDTRL NKAVWAKGIRNVPYRIRVRLSRKRNEDEDSP NKLYTLVTYVPVTTFKNLQTVNVDEN
1017	2367	A	8513	54	1196	LERTPASADMAWTKYQLFLAGLMLVTGSINT LSAKWADNFMAEGCGGSKEHSFQHPFLQAV GMFLGEFSCLAAFYLLRCRAAGQSDSSVDPQ QPFNPLLFLPPALCDMTGTSLMYVALNMTSA SSFQMLRGAVIIFTGLFSVAFLGRRLVLSQWL GILATIAGLVVVGLADLLSKHDSQHKLSEVIT GDLLIIMAQIIVAIQMVLEEKFVYKHNVHPLR AVGTEGLFGFVILSLLLVPMYYIPAGSFSGNP RGTLEDALDAFCQVGQQPLIAVALLGNISSIA FFNFAGISVTKELSATTRMVLDSLRTVVIWAL

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1018	2368	A	8518	324	694	SPFWTEKRRMEKPLFPLVPLHWFGFGYTALV VSGGIVGYVKTGSVPSLAAGLLFGSLAGLGA YQLYQDPRNVWGFLAATSVTFVGVMGMRS YYYGKFMPVGLIAGASLLMAAKVGVRMILM TSD
1019	2369	A	8526	2	1787	VSAAAVNMEPPDAPAQARGAPRLLLLAVLL AAHPDAQAEVRLSVPPLVEVMRGKSVILDCT PTGTHDHYMLEWFLTDRSGARPRLASAEMQ GSELQVTMHDTRGRSPPYQLDSQGRLVLAEA QVGDERDYVCVVRAGAAGTAEAAARLNVF AKPEATEVSPNKGTLSVMEDSAQEIATSNSRN GNPAPKITWYRNGQRLEVPVEMNPEGYMTS RTVREASGLLSLTSTLYLRLRKDDRDASFHC AAHYSLPEGRHGRLDSPTFHLTLHYPTEHVQ FWVGSPSTPAGWVREGDTVQLLCRGDGSPSP EYTLFRLQDEQEEVLNVNLEGNLTLEGVTRG QSGTYGCRVEDYDAADDVQLSKTLELRVAY LDPLELSEGKVLSLPLNSRAVVNCSVHGLPTP ALRWTKDSTPLGDGPMLSLSSITFDSNGTYVC EASLPTVPVLSRTQNFTLLVQGSPELKTAEIEP KADGSWREGDEVTLICSARGHPDPKLSWSQL GGSPAEPIPGRQGWVSSSLTLKVTSALSRDGI SCEASNPHGNKRHVFHFGTVSPQTSQAGVAV MAVAVSVGLLLLVVAVFYCVRRKGGPCCRQ RREKGAP
1020	2370	A	8530	2	1200	PRVRLLRPSRSRSCRGLLSTRAPGPSPFRSLHS SPLLPHAMKSPFYRCQNTTSVEKGNSAVMGG VLFSTGLLGNLLALGLLARSGLGWCSRRPLR PLPSVFYMLVCGLTVTDLLGKCLLSPVVLAA YAQNRSLRVLAPALDNSLCQAFAFFMSFFGL SSTLQLLAMALECWLSLGHPFFYRRHITLRLG ALVAPVVSAFSLAFCALPFMGFGKFVQYCPG TWCFIQMVHEEGSLSVLGYSVLYSSLMALLV LATVLCNLGAMRNLYAMHRRLQRHPRSCTR DCAEPRADGREASPQPLEELDHLLLALMTV LFTMCSLPVIYRAYYGAFKDVKEKNRTSEEA EDLRALRFLSVISIVDPWIFIIFRSPVFRIFFHKI FIRPLRYRSRCSNSTNMESSL
1021	2371	A	8536	1	237	RRGEIDMATEGDVELELETETSGPERPPEKPR KHDSGAADLERVTDYAEEKEIQSSNLETAMS VIGDRRSREQKAKQER
1022	2372	A	8537	94	541	RKERRRRRMEAVVFVFSLLDCCALIFLSV YFIITLSDLECDYINARSCCSKLNKWVIPELIG HTIVTVLLLMSLHWFIFLLNLPVATWNIYRYI MVPSGNMGVFDPTEIHNRGQLKSHMKEAMI KLGFHLLCFFMYLYSMILALIND
1023	2373	A	8540	26	431	RMMKCPQALLAIFWLLLSWVSSEDKVVQSPL SLVVHEGDTVTLNCSYEVTNFRSLLWYKQEK KAPTFLFMLTSSGIEKKSGRLSSILDKKELSSIL NITATQTGDSAIYLCAVEAQCSLVTCSLYSNS TAEALQL
1024	2374	A	8544	1731	743	GVRLRYSPIAVVMVGEAGRDLRRRRAVAVT AEKMAVLAPLIALVYSVPRLSRWLAQPYYLL SALLSAAFLLVRKLPPLCHGLPTQREDGNPCD FDWREVEILMFLSAIVMMKNRRSITVEQHIGN IFMFSKVANTILFFRLDIRMGLLYITLCIVFLM

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1025	2375	A	8546	2194	1707	VIREFNLNELYQRAKKLSKAGDNIPEEQPVAS TPTTVSDGENKKDK TVSFHKTMASLKCSTVVCVICLEKPKYRCPA
1023	2373		0540	2131		CRVPYCSVVCFRKHKEQCNPETRPVEKKIRS ALPTKTVKPVENKDDDDSIADFLNSDEEEDR VSLQNLKNLGESATLRSLLLNPHLRQLMVNL DQGEDKAKLMRAYMQEPLFVEFADCCLGIV EPSQNEES
1026	2376	A	8547	1078	594	VGMELPAVNLKVILLGHWLLTTWGCIVFSGS YAWANFTILALGVWAVAQRDSIDAISMFLGG LLATIFLDIVHISIFYPRVSLTDTGRFGVGMAIL SLLLKPLSCCFVYHMYRERGGELLVHTGFLG SSQDRSAYQTIDSAEAPADPFAVPEGRSQDAR GY
1027	2377	A	8557	1	340	DFLGPASPQEEGGSESSTMTELETAMGMIIDV FSRYSGSEGSTQTLTKGELKVLMEKELPGFLQ SGKDKDAVDKLLKDLDANGDAQVDFSEFIVF VAAITSACHKYFEKAGLK
1028	2378	A	8569	20	963	KMAATLGPLGSWQQWRRCLSARDGSRRLLL LLLLGSGQGPQQVGAGQTFEYLKREHSLSKP YQGEAPRPCFLRDWELQVHFKIHGQGKKNL HGDGLAIWYTKDRMQPGPVFGNMDKFVGLG VFVDTYPNEEKQQERVFFYISAMVNNGSLSY DHERDGRPTELGGCTAIVRNLHYDTFLVIRY VKRHLTIMMDIDGKHEWRDCIEVPGVRLPRG YYFGTSSITGDLSDNHDVISLKLFELTVERTPE EEKLHRDVFLPSVDNMKLPEMTAPLPPLSGL ALFLIVFFSLVFSVFAIVIGIILYNKWQEQSRK RFY
1029	2379	A	8572	İ	578	AAAASHRSRARSRPRRVSSGPAPRRAQSSAG RVASGLDSAPLCTMARALCRLPRRGLWLLLA HHLFMTTACQEANYGALLRELCLTQFQVDM EAVGETLWCDWGRTIRSYRELADCTWHMAE KLGCFWPNAEVDRFFLAVHGRYFRSCPISGR AVRDPPGSILYPFIVVPITVTLLVTALVVWQS KRTEGIV
1030	2380	A	8574	1352	372	DSSTVKGGSESRHLCLIPDLKGKARTREASSG SRTCGRRTSLCTSAKSSWTYRSGRLSWQSIKG THLTITQALRQPLHRAPLLPGQLCWSPRPLEK NKAMGRPLLLPLLLLLQPPAFLQPGGSTGSGP SYLYGVTQPKHLSASMGGSVEIPFSFYYPWEL AIVPNVRISWRRGHFHGQSFYSTRPPSIHKDY VNRLFLNWTEGQESGFLRISNLRKEDQSVYF CRVELDTRRSGRQQLQSIKGTKLTITQAVTTT TTWRPSSTTTIAGLRVTESKGHSESWHLSLDT AIRVALAVAVLKTVILGLLCLLLLWWRRRKG SRAPSSDF
1031	2381	A	8580	905	340	RRTAGIYPCFPKPGRTRHALCSVVLLLLTGQL AFDDFQESCAMMWQKYAGSRRSMPLGARIL FHGVFYAGGFAIVYYLIQKFHSRALYYKLAV EQLQSHPEAQEALGPPLNIHYLKLIDRENFVDI VDAKLKIPVSGSKSEGLLYVHSSRGGPFQRW HLDEVFLELKDGQQIPVFKLSGENGDEVKKE
1032	2382	A	8593	2558	961	RRRPRLLPGAEPCEPRVGPRRADMGCSAKAR WAAGALGVAGLLCAVLGAVMIVMVPSLIKQ

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1033	2383		9505	595	,	ESGAMEGETLHTFYTQLVLMPKVMHYAQYV LLALGCVLLLVPVICQIRSQEKCYLFWSSSKK GSKDKEAIQAYSESLMTSAPKGSVLQEAKL
1033	2383	A	8595	393	767	AHLPDTLLLPPHSPTVPTPKSFQCSQKACFSRS FCLLLSLVSSSLVSLSLCPPLTQA
1034	2384	A	8597	640	164	VTTSCIIPFAFGLGVRASERLAEIDMPYLLKYQ PMMQTIGQKYCMDPAVIAGVLSRKSPGDKIL VNMGDRTSMVQDPGSQAPTSWISESQVFQTT EVLTTRITELQRRFPTWTPDQYLRGGLCAYSG GAGYVRSSQDLSCDFCNDVLARAKYLKRHG F
1035	2385	A	8603	936	204	AMASTLEYSPSPLRRLVGPAAGFSRAARADL SWDPMAFFTGLWGPFTCVSRVLSHHCFSTTG SLSAIQKMTRVRVVDNSALGNSPYHRAPRCI HVYKKNGVGKVGDQILLAIKGQKKKALIVG HCMPGPRMTPRFDSNNVVLIEDNGNPVGTRI KTPIPTSLRKREGEYSKVLAIAQNFV
1036	2386	A	8606	1	562	PTRAHSFDLCCSPCRRRLLGREEAGEEPTSPV TQYLQPRSPEECKMFACAKLACTPSLIRAGSR VAYRPISASVLSRPEASRTGEGSTVFNGAQNG VSQLIQREFQTSAISRDIDTAAKFIGAGAATVG VAGSGAGIGTVFGSLIIGYARNPSLKQQLFSY AILGFALSEAMGLFCLMVAFLILFAM
1037	2387	A	8615			SPGPSLPESAESLDGSQEDKPRGSCAEPTFTDT GMVAHINNSRLKAKGVGQHDNAQNFGNQSF EELRAACLRKGELFEDPLFPAEPSSLGFKDLG PNSKNVQNISWQRPKDIINNPLFIMDGISPTDI CQGILGDCWLLAAIGSLTTCPKLLYRVVPRG QSFKKNYAGIFHFQIWQFGQWVNVVVDDRL PTKNDKLVFVHSTERSEFWSALLEKAYAKLS GSYEALSGGSTMEGLEDFTGGVAQSFQLQRP PQNLLRLLRKAVERSSLMGCSIEVTSDSELES MTDKMLVRGHAYSVTGLQDVHYRGKMETLI RVRNPWGRIEWNGAWSDSAREWEEVASDIQ MQLLHKTEDGEFWMSYQDFLNNFTLLEICNL TPDTLSGDYKSYWHTTFYEGSWRTGSSAGGC RNHPGTFWTNPQFKISLPEGDDPEDDAEGNV VVCTCLVALMQKNWRHARQQGAQLQTIGFV LYAVPKEFQNIQDVHLKKEFFTKYQDHGFSEI FTNSREVSSQLRLPPGEYIIIPSTFEPHRDADFL LRVFTEKHSESWELDEVNYAEQLQEEKVSED DMDQDFLHLFKIVAGEGKEIGVYELQRLLNR MAIKFKSFKTKGFGLDACRCMINLMDKDGSG KLGLLEFKILWKKLKKWMDIFRECDQDHSGT LNSYEMRLVIEKAGIKLNNKVMQVLVARYA DDDLIIDFDSFISCFLRLKTMFTFFLTMDPKNT GHICLSLEQVLGEGWEGICRIAPACPSTPPPPS

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion SDVPGPASCPRLFPPWDLLPVSTVAADDHVGI EAL
1038	2388	A	8621	3	1494	RSRMARAPLGVLLLLGLLGRGVGKNEELRLY HHLFNNYDPGSRPVREPEDTVTISLKVTLTNL ISLNEKEETLTTSVWIGIDWQDYRLNYSKDDF GGIETLRVPSELVWLPEIVLENNIDGQFGVAY DANVLVYEGGSVTWLPPAIYRSVCAVEVTYF PFDWQNCSLIFRSQTYNAEEVEFTFAVDNDG KTINKIDIDTEAYTENGEWAIDFCPGVIRRHH GGATDGPGETDVIYSLIIRRKPLFYVINIIVPCV LISGLVLLAYFLPAQAGGKCTVSINVLLAQT VFLFLIAQKIPETSLSVPLLGRFLIFVMVVATLI VMNCVIVLNVSQRTPTTHAMSPRLRHVLLEL LPRLLGSPPPPEAPRAASPPRRASSVGLLLRAE ELILKKPRSELVFEGQRHRQGTWTAAFCQSL GAAAPEVRCCVDAVNFVAESTRDQEATGEE VSDWVRMGNALDNICFWAALVLFSVGSSLIF LGAYFNRVPDLPYAPCIQP
1039	2389	A	8636	1	900	PGRERPGGGGARRRPQHLPALLPSERPDCATL QAMENELPVPHTSSSACATSSTSGASSSSGCN NSSSGGSGRPTGPQISVYSGIPDRQTVQVIQQ ALHRQPSTAAQYLQQMYAAQQQHLMLQTA ALQQHLSSAQLQSLAAVQQASLVSNRQGST SGSNVSAQAPAQSSSINLAASPAAAQLLNRA QSVNSAAASGIAQQAVLLGNTSSPALTASQA QMYLRAQMLIFTPTATVATVQPELGTGSPAR PPTPAQVQNLTLRTQQTPAAAASGPTPTQPVL PSLALKPTPGGSQPLPTPA
1040	2390	A	8645	98	1388	ASQLAFGGKLTSTPSRDFQGCGRGAVTCCSF HEHRHQSGRCLSTGMAPNLKGRPRKKKPCPQ RRDSFSGVKDSNNNSDGKAVAKVKCEARSA LTKPKNNHNCKKVSNEEKPKVAIGEECRADE QAFLVALYKYMKERKTPIERIPYLGFKQINLW TMFQAAQKLGGYETITARRQWKHIYDELGG NPGSTSAATCTRRHYERLILPYERFIKGEEDKP LPPIKPRKQENSSQENENKTKVSGTKRIKHEIP KSKKEKENAPKPQDAAEVSSEQEKEQETLISQ KSIPEPLPAADMKKKIEGYQEFSAKPLASRVD PEKDNETDQGSNSEKVAEEAGEKGPTPPLPSA PLAPEKDSALVPGASKQPLTSPSALVDSKQES KLCCFTESPESEPQEASFPRLPHHTGHRWQTR MRRRMTNCPPWQITLPTAP
1041	2391	A	8646	538	1492	LLQEMCTKTIPVLWGCFLLWNLYVSSSQTIYP GIKARITQRALDYGVQAGMKMIEQMLKEKK LPDLSGSESLEFLKVDYVNYNFSNIKISAFSFP NTSLAFVPGVGIKALTNHGTANISTDWGFESP LFVLYNSFAEPMEKPILKNLNEMLCPIIASEVK ALNANLSTLEVLTKIDNYTLLDYSLISSPEITE NYLDLNLKGVFYPLENLTDPPFSPVPFVLPER SNSMLYIGIAEYFFKSASFAHFTAGVFNVTLS TEEISNHFVQNSQGLGNVLSRIAEIYILSQPFM VRIMATEPPIINLQPGNFTLDIPASIMMLTQPK NSTVETIVSMDFVASTSVGLVILGQRLVCSLS LNRFRLALPESNRSNIEVLRFENILSSILHFGVL PLANAKLQQGFPLPNPHKFLFVNSDIEVLEGF LLISTDLKYETSSKQQPSFHVWEGLNLISRQW RGKSAP ARRIARTRESKAAVSQDNVPALQPGKKKKLR
		**	55,2			LGGKKKKFKFFRLPKEFKKQLMYSPSNFKKM

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Scrine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  TSLAGNTVQCLNKLKYVIYSAQYPAYGNITT LDMITSTDHVLEQDFWICFTFYSVKERQI GLKTRAPATPTFQREVLGPAKQDMQRRCPRI GLMTSLIKPIKRRWRDYKRWKSGGFTGESC
			ļ			HHADTLGDRGGLQGDHSELLQWQKRILRTE GEPSPKYISKNIFPICSYITGFL
1044	2394	Ā	8718	292	1490	GTVKTSVATPITAGHSCSSGGVLQVKSPATQS GFKFTSKMEDFNMESDSFEDFWKGEDLSNYS YSSTLPPFLLDAAPCEPESLEINKYFVVIIYAL VFLLSLLGNSLVMLVILYSRVGRSVTDVYLL NLALADLLFALTLPIWAASKVNGWIFGTFLC KVVSLLKEVNFYSGILLLACISVDRYLAIVHA TRTLTQKRYLVKFICLSIWGLSLLLALPVLLFR RTVYSSNVSPACYEDMGNNTANWRMLLRIL PQSFGFIVPLLIMLFCYGFTLRTLFKAHMGQK HRAMRVIFAVVLIFLLCWLPYNLVLLADTLM RTQVIQETCERRNHIDRALDATEILGILHSCLN PLIYAFIGQKFRHGLLKILAIHGLISKDSLPKDS RPSFVGSSSGHTSTTL
1045	2395		8724	254	3184	FRANLAITVANRRGAQGGKMHTCCPPVTLEQ DLHRKMHSWMLQTLAFAVTSLVLSCAETIDY YGEICDNACPCEEKDGILTVSCENRGIISLSEIS PPRFPIYHLLLSGNLLNRLYPNEFVNYTGASIL HLGSNVIQDIETGAFHGLRGLRRLHLNNNKL ELLRDDTFLGLENLEYLQVDYNYISVIEPNAF GKLHLLQVLILNDNLLSSLPNNLFRFVPLTHL DLRGNRLKLLPYVGLLQHMDKVVELQLEEN PWNCSCELISLKDWLDSISYSALVGDVVCETP FRLHGRDLDEVSKQELCPRRLISDYEMRPQTP LSTTGYLHTTPASVNSVATSSSAVYKPPLKPP KGTRQPNKPRVRPTSRQPSKDLGYSNYGPSIA YQTKSPVPLECPTACSCNLQISDLGLNVNCQE RKIESIAELQPKPYNPKKMYLTENYIAVVRRT DLLEATGLDLLHLGNNRISMIQDRAFGDLTN LRRLYLNGNRIERLSPELFYGLQSLQYLFLQY NLIREIQSGTFDPVPNLQLLFLNNNLLQAMPS GVFSGLTLLRLNLRSNHFTSLPVSGVLDQLKS LIQIDLHDNPWDCTCDIVGMKLWVEQLKVG VLVDEVICKAPKKFAETDMRSIKSELLCPDYS DVVVSTPTPSSIQVPARTSAVTPAVRLNSTGA PASLGAGGGASSVPLSVLILSLLLVFIMSVFVA AGLFVLVMKRRKKNQSDHTSTNNSDVSSFN MQYSVYGGGGGTGGHPHAHVHHRGPALPK VKTPAGHVYEYIPHPLGHMCKNPIYRSREGN SVEDYKDLHELKVTYSSNHHLQQQQPPPPP QQPQQPPPQLQLQPGEEERRESHHLRSPAYS VSTIEPREDLLSPVQDADRFYRGILEPDKHCST TPAGNSLPEYPKFPCSPAAYTFSPNYDLRRPH QYLHPGAGDSRLREPVLYSPPSAVFVEPNRNE YLELKAKLNVEPDYLEVLEKQTTFSQF
1046	2396	A	8736	28	452	SPSAAGGLAWVSLALGSGSRGRDHSGSGVGT AMAGALVRKAADYVRSKDFRDYLMSTHFW GPVANWGLPIAAINDMKKSPEIISGRMTFALC CYSLTFMRFAYKVQPRNWLLFACHATNEVA QLIQGGRLIKHEMTKTASA
1047	2397	A	8741	673	924	ALPGTPQQTVTLNTDGKVKSFTSPHSNPNLPP AKFFTSLQSLNWSSHLPPSPATESVGKRGNAK
1048	2398	A	8747	3	5054	PPTTKLLHSSPLWNFFAQQL PEVTKPSLSQPTAASPIGSSPSPPVNGGNNAKR

SEQ ID SEQ ID NO. of nucleotide of peptide collide sequence (A-alianine C-Cysteine, 1944) but one of peptide social sequence (A-alianine C-Cysteine, 1944) beautiful in succession of peptide social sequence (A-alianine C-Cysteine, 1944) beautiful in succession of peptide social sequence (A-alianine, C-Cysteine, 1944) beautiful in succession of peptide social residue of peptide soc	CODE TO	CEC VE		CEC			
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residue of peptide sequence	uence			914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
peptide   /-possible nucleotide deletion,  -possible nucleotide insertion		ļ			amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
peptide   /-possible nucleotide deletion,  -possible nucleotide insertion				1	residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
sequence   nucleotide insertion	ļ	}	Į		peptide		
VAVPNGOPPSAARYMREVPERRECOGDEN' VILKERQOPPPSCMLIGGGAGPPCTARGAN PRINAQVITGALLQESGTAPDSTLGGAAASIN PRINAQVITGALLQESGTAPDSTLGGAASIN ANSTWOSGASSNINGTSPIPHIMDKVIVDG'S DMEEWPCLASKDTESSSENTIDNINSASNPGGE KSTLFGSTTISNKGKGSQCGASSGEALGV WKSDPFARKSVQSSNSTITENNINGLGNWRYN'S GORIGGGGGFSHAPPNINSRS APAL VQEGI'S RKGALETIDNINSASQVSTVGGTSREQQSKME  NAGVNPVVSGREQAQHINTOPFRINGINSE. NLSSPNPMENKGMPFGMGIGNTSRSTDAPSG STGDRKTGSVGSWGAARGPSGTDTVSGGSNS GNNGNNGKEREDSWKGASVQKSTGSKNDS WDNNINSTIGGSWNFGPQDSNDINKWEGDN MTSGVSQGEWKQPTGSDELKIGEWSQFNOON STGAWDNQKGHPLENQGNAQAPCWGRSS STGSEVEGGSTGSNHKAGSSDSHNSGRSSY RTHHPDCQAVLQTLLSRTDLDPFVSNTGWG QTQIKQDTVWDIEEVPRFSGKSDKGTEGWSS RTHYDDCQAVLQTLLSRTDLDPFVSNTGWG QTQIKQDTVWDIEEVPRFSGKSDKGTEGWSS AATQTKNSGGWGDAPSCSNGWKSGWGELS AATTEWKDPKNTGGWDYKNNINSSNWGGGR PDEKTPSSWWEIPSSKDQCWGGGGPQNGWS SGKNGWGEEVDQTTRNSWSSASKSVGCGS PERSSWEDSSSCOORDSWSGAGGWEDCKS AATGWNGGGNASLASKGGWEDCKS PAWNETGRQPNSWNKQHQQQPPQOGPPPQ PASAGSWGGPPPPGON STRSSWSGFQPA TRYDERSGWEFSSPQSISRKMDIDDGTSAWG DPGSTYNTNINALWKNNSQGQPRGPAGWS WKSGAGGSWGGPPGON STRYNSWSAGGWEDCKS PAWNETGRQPNSWNKQHQQQPPQOGPPPQ PASAGSWGGPPPFQNVRINSNSWSSGFQPA TRYDERSGWEFSSPQSISRKMDIDDGTSAWG DPGSTYNTNINALWKNNSQGGPAFRALLTP MTSKASJSKSMDDGWGESDGYTTGARHEPS WKSGAGGSWGTHASSGCASKSTOON AMANGGPPDBMRKDRGGGFRONGTTDA MCGCSWGGPPTGANGSWGGGGGGAGAAAAAAAAAAAAAAAAAAAAAAAAAA							
VILIKBGOPPPSCMLLGGGAGPPPCTAPGAG PNAQVTGALLQSEGATAPDSTLGGAAASNY ANSTWGSGASSNNGTSPIPHIWDKVIVDGS DMEEWCLASKDTESSSENTITDNSASPGGS KSTLPGSTTSNKGKGSQCQSASSGNECNLGV WKSDPFAKSVQSSNSTTRINNGCMWRVS GQDRIGPGSSFSNTFIPNNSTSAWPALVQGGTS RGAGLAETDNSNSSAQVSTVGGTSCROWKNVS GQDRIGPGSSFSNTFIPNNSTSAWPALVQGGTS RGAGLAETDNSNSSAQVSTVGGTSCROSKME NAGVNFVVSGREQAQHINTDGPKNGNTNSL NLSSFNMENKGMPFGMGLGNTSSTDAFSQ STORKTGSVGSWGAARGPSGTDTVSGGSSK STORKTGSVGSWGARGPSGTDTVSGGSSK GNNGNKCKEREDSWKGASVQKSTGSKNDS WDNNNRSTGGSWNFGPQDSNDKWGEGRNQ MTSGSVGGWKGPTGSDELKIGEWSGPNQPN SSTGAWDNQKGFPLLENVGGNAQAPCWGRSS STGSEVEGQSTGSNHKAGSSDSHNSGGRSV QTQIKGDTVWDIEGWSGPNQPN SSTGAWDNQKGFPLLENVGGNAQAPCWGRSS STGGEVFQGSTGSNHKAGSSDSHNSGRSSV RPTHPDCQAVLOTLLSRTDLDPPLVLSNTGWG QTQIKGDTVWDIEGWSGPNQMKSGWGELS SATGWEDPROFTSSENGEGSWGFTSSKNGRSS SGROWGEVDTLTSNSWESSASSWGGRS PDEKTPSSWAENPSKDQCWGGGRQPNQOW SGROWGEVDTLTNSNWESSASKEVSGWG GGQNEIGTWGNGGNASLASKGGWBCLKS SGROWGESWDTTNSNWESSASKEVSGWG EGGQNEIGTWGNGGNASLASKGGWBCKSK SGROWGESWDGTNSNWESSASKEVSGWG GGGNEIGWGGGROPNOWN PROPER PASAGSWGGPPTPPFGNVRSNSSWSGPPPPP PASAGSWGOPPTPFFGNVRSNSSWSGGPQPP PASAGSWGGPPTGTPFGNVRSNSSSSSGNGGG GKKQMGEVDGGGGRDDSWNNPLAKGPSNDG LISQTEDNFSIKMLSVGSLSDKKFDVDKA MEEEBDGGVWTTTGSQGSASSINSASWGG GKKQMCSLKGGNDDSWNNPLAKGPSNDG LISQTEDNFSIKMLSVGSLSDKKFDVDKA MEEEBDGGVWTTTGSQGSASSINSASWGG GKKGKLKGRNDSWNNPLAKGPSNDG LISQTEDNFSIKMLSVGSLSDKKFDVDKA MSALQQQQQQCQRGRGFRGFRGMF WEEEBDGGVWTTGSQGGASSINGSSWGGG GKKQMCSLKGRNDSWNNPLAKGPSPND GRIPGPTGLFSQVASMAKRGNGFPTSSLFVG MEEEBDGGVWTTGSQGGASSINGSSWGGG GKKQMCSLKGRNDSWNPLAKGPSPND GRIPGPTGLFSQVASMAKRGNGFTSNFVGFTS GPTFEKGGSIGLFGNSTAQSRGLHTTVOPLN SPSLRAQVPPTGATTSPIVDTDHQLLRDNTTGS GPTFEKGGSIGLFGNSTAQSRGHPTSSFK MTTLTHPTPGTLTNFTSSFWSTAFSNFW NGPDIFGDTLGGTTGPAGDSWLPAKSPSPNV LILLICASTSLUGAGGSGVRFSVWLVILLINTGRS NSSLNTSLEPFGAWPYSASNSSFTNVHSTSAK FPDXKSTWSDPFIGNFTHLSNKMWKNHISS RNTTLTPRPPGGNTGLGGSTSG SGGMDTGGGSNSSSACTLDSFFPFIAPWG SSCALAGASASTWSDGGSNRSVWLVILLILIS TAMYGALAACHTALCHUDICAGTTILLLS TAMYGALAACHTALCHUDICAGTTILLLS TAMYGAHAPLLALCHUDICAGTSPFFAPWGA		· · · · · · · · · · · · · · · · · · ·	<del>                                     </del>		Doquesion		
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ANSTWOSGASSNNIGTSPNPHHWDKYIVDGS DMEEWPCLASKDTESSSENTIDNNSASNPOSE KSTLPGSTTSNKOKGSQCQSASSGNECALGV WKSDPKAKSVQSSNSTTENNNIGLGNWRNVS GQDRIGPGSGESSPNPINSNIGLGNWRNVS GQDRIGPGSGESSPNPINSNIGLGNWRNVS GQDRIGPGSGESSPNPINSNIGLGNWRNVS GQDRIGPGSGESSPNPINSNIGLGNWRNVS GRINGSTORT AND ANG ANG ANG ANG ANG ANG ANG ANG ANG ANG	1	ļ	ł	}		}	
DMEEWPCLASKOTESSSENTTONNASHOENER  KSTLPGSTTISKKGKGSGOCSASSGOBECNLGV WKSDPKAKSVOSSNSTTENNNGLGNWRNVS GORDIGFGSGFSINPINNSHAPALVOGGTS RKGALETDNINSSAQVSTVGGTSREQQSISME NAGWIFVVSGREQQINTORFINGNITNSI. NLSSPPMENKGMPFGMGLGNTSRSTIDAPQ STGDRKTGSVGSWGAARAFSCHGNGNTNSI. NLSSPPMENKGMPFGMGLGNTSRSTIDAPQ STGDRKTGSVGSWGAARAFSCHJVSGTSKANDS WDNNRSTGGSWHGPGDSDNDKWGEGNK MTSGVSOGEWKQPTGSDELKIGEWSGPNOPN SSTGAWDNOKGHLLENOGMAQAPCWGRSS STGSEVEGQSTGSNHKAGSSDSINNSGRSV RPTHPDCQAVLOTILSRTDLWSDNOW GTQIKGDDTVWDIEEVPPPGKSDKGTEGWES AATOTKNSGGMGAPSSGOMKGSWGERSV RPTHPDCQAVLOTILSRTDLWSDWGGR PDEKTPSSWNENPSKDGCWGGROPNOGWS SGKNGWGEUDQTKNNWSSASKPVSGWG GGGONEIGTWGROGANSWSGGRPNOGWS SGKNGWGEUDQTKNNWSSASKPVSGWG GGGONEIGTWGROGANSASKPVSGWG GEGONEIGTWGNOWSSASKPVSGWG GGGONEIGTWGNOWSSASKPVSGWG GGGNEGWGPSSWNKOPLAKAGS AATOTKNSGGGAPPREPRILTPT PATSKSASDSKSMQDGGGPPPPPPP PLASGSWGGGPPPPPPPNSWNSSSGPQPA TPEDEEPSGWEEPSPGSISKKMDIDDOTAAWG DPNSYNYXNNUNUNNSQGGPAPREPRILTPT PMTSKSASDSKSMQDGWGESDGPVTGARHPS WEEEEDGGVWNTTGSOGGAPGPREPRILTPT ANTICKSASDSKSMQDGWGESDGPVTGARHPS WEEEEDGGVWNTTGSOGGAPGPREPRILTPT ANTICKSASDSKSMQDGWGESDGPVTGARHPS WEEEEDGGVWNTTGSOGGAPGREPRILTPT ANTICKSASDSKSMQDGWGESDGPVTGARHPS WEEEEDGGVWNTTGSOGGAPGREPRILTPT ATTICKSASDSKSMQDGWGESDGPVTGARHPS WEEEEDGGVWNTTGSOGGAPGREPRILTPT ANTICKSASDSKSMQDGWGESDGPVTGARHPS WEEEEDGGVWNTTGSOGGAPGREPRILTPT ANTICKSASDSKSMQDGWGESDGPVTGARHPS WEEEEDGGVWNTTGSOGGAPGRAFTISS GGKKGMKGLKGLKGGNNDSWNTPLAKOGFTPPS WEEEEDGGVWNTTGSOGGAPGRAFTISS GGKKGMKGLKGLKGGNNDSWNTPLAKOGFTPPSCLSP GRANGWGESDGPVTGARHPS WEEEEDGGVWNTTGSOGGAPGRAFTISS GGRADLAGALVGGPTPRONGCACG LLQQQQQQCLQNQKKISSTFYOYR PHILDNWPPALAVGLPDLJTGGPTGVGGGF SGGGADLAGALVGPTWGGNDDPSDP YVTPGSVLGGTATSPIVDTDGQLLRDNTTGS NSSLAVTLSPSFPTGTGVWRKGIONDPFSDP YVTPGSVLGGTATSPIVDTDGQLLRDNTTGS NSSLAVTLSPSFPTGVGTWWRGKACHTENTUTGGT LRTSTKGQAATSPIVDTDGQLLRDNTTGS NSSLAVTLSPSFPTGVGTWWRGKACHTENTUTGTA LRTSTKGQAATSPIVDTDGQCGCQLACACACACACACACACACACACACACACACACACACA						{	
KSTLPGSTTSNKGKGSQCQSASSGNECNLGV WKSDPKAKSVQSSNSTTENGLGNWRNVS GQDRIGPGGGESNFRNNSRPSAWPALVQEGTS RKGALETDNSNSAQVSTVQCFGCYSEQGKME RKGALETDNSNSAQVSTVQCFGCYSEQGKME RKGALETDNSNSAQVSTVQCSGCYSEQGKME RKGALETDNSNSAQVSTVQCSGCYSEQGKME RKGALETDNSNSAQVSTVQCSGNS GNNGNMGEREDSWKGASVSTGSNTSSTDAPSQ STODRXTGSVGSWGAARGPSGTDTVSGQSNS GNNGNMGEREDSWKGASVSTGSNTSS GNNGNMGEREDSWKGASVSTGSNTS WDNNRRSTGGSWNRGPQDSNDMKWGBONK MTSGVSQGEWKOPTGSDELKIGEWGSPNQPN SSTGAWDNQKGHPLLENQGMAQAPCWGRSS SSTGSEVEGQSTGSMIKAGSSDSINSGRSY RPIHPDCQAVLQTILSRTGLDIPRVLSNTGWG QTQTKQDTVWDIEEVPRPSGKSDKGTGEWS AAATQTKNSGGMGAPSQSMKKSGWGELS ASTEWKDPKNTGGWNDYKNNNSSNGGR PDEKTPSSWNENPSKDQWGGGRQPNQGWS SGKNGWGEWDQTKNNNWESSASKPVSGWG EGGQNEIGTWGNGGASLASKGWEDCKRS, PAWSETGRQPNSWNKOHQQQQPPQQPPPQ PEASGSWGGPTPPPFONNVRSSNSSWGPQCAR PREMEPSGWEEPSPQSISKKMDIDDGTSAWG DPNSYNYKNVN,WDKNSQGGAPPREPNLTPT MTSKSASDSKSMQDGWGESDGPVTGARHPS WEEEEDGGWWTTGNGGGASSHNSASWGQG GKKQMKCSLKGGNNDSWMNPLAKGPSNMG LLSQTEDNPSSKMDLSOGGAPREPNLLTPT MTSKSASDSKSMQDGWGSSDGPVTGARHPS WEEEEDGGWWTTGNGGGASSHNSASWGQG GKKQMKCSLKGGNNDSWMNPLAKGPSNMG LLSQTEDNPSSKMDLSVGSTARHPS WEEEEDGGWWTTGNGGGASSHNSASWGQG GKKQMKCSLKGGNNDSWMNPLAKGPTNSGLSP GLFNVGPQLSPQQQQQQLLONQKRISQAYRQCGGCQDA RMNLGDFNDIMRLDRSGFRPPNSKDMGTTDS GPYFEKGGSHGLFGNSTAGSGGHTPVQPLN SSPSLRAQVPPQTSPQVSASMLKQFFTNSGLSP GLFNVGPQLSPQQQQQLLONQKRISQAYRQCGGPQA RMVSALQQQQQQQLLONQKRISQAYRQCGGPA RMVSALQQQQQQQQCQCGGGGRKGMSFSSPVGFK RMSALQQQQQQQCQCGGGGRKGMSFSSPVGFK RMSALQQQQQQQCQCGGGGRAFTSSRVTDLLADNTTGS NSSLNTSLPSFGAWPYSASDNSTTNVTISAK FPDYKSTWSPPGGAPRSAMJKNCHHSS RNTTPLPRPPGGLTNFKPSSFWSTAPESVG WGTQDSELASASTWSDGGSYSSM (MGSSSA GGSSGADLAGALAQQPTPAALTSAMWXNHISS RNTTPLPRPPGGLTNFKPSSFWWSTAPESVG WGTQDSELASASTWSDGGSYSSM (MGSSSA GGSSGADLAGALAQQPTPAALTSAMWXNHISS RNTTPLPRPPGGLTNFKPSSFWWSTAPESVG WGTQDSELASASTWSDGGSYSSM (MGSSSA GGSSGADLAGALAQQPTPAALTSAMWXNHILS RNTTLEQROPHTLALTCHLARWTT.MLLS TAMYGABAPLLALCHVOGNVPFRPSSALVLT LETKILLCAFSTGLOWGGFPVPRQSAALLT LETKILLCAFSTGLOWGGFPPVRQAA							
WKSDPKAKSVQSSNSTTENNNGLGNWRNVS GQDRIGPGSGFSNPNNNSNSAPALVQGGTS RKGALETDNSNSAQVSTVQGTSREQQSKME NAGWNFVVSGREQQIHNTDSFNGNTNSL NLSSPPMENKGMPFOMGLGNTSRSTDAPQ STGDRKTGSVGSWGAARAPGSTDTVSQGSNS STGDRKTGSVGSWGAARAPGSTDTVSQGSNS GNNGNNGKERDSWKGASVQKSTGSKNDS WDNNRNSTGSWNFGPQDSDNDKWGEGNK MTSGVSQGEWKQPTGSDELKIGEWSGPNOPN SSTGAWDNQKGHPLLENQGNAQAPCWGRSS STGSEVEGQSTGSNHKAGSSDSHNSGRSV RPTHPDCQAVLOTILLSRTDAPSLSNTGWG QTQIKGDTVWDIEEVPRPEGKSDKGTEGWES AATOTKNSGGWGAPSSCONKGSWGESY RPTHPDCQAVLOTILLSRTDAWSGWGESY RPTHPDCQAVLOTILLSRTDAWSGWGESY SGKNGWGEWDOWNSCSWGESS SGKNGWGEWDOWNSCSWGESS AATOTKNSGGWGAPSSCONKGSWGESY RPDEKTPSSWNSHPSKDGCWGROPNOGWS SGKNGWGEDDOWNTKNNSSNSWGGOP PEEKTPSSWNSHPSKDGCWGDPVGARS SGKNGWGEDDOWNTSNSWSSASKPVSGWG EGGONEIGTWGNOGNASASKPVSGWG EGGONEIGTWGNOGNASASKPVSGWG EGGONEIGTWGNNGWGSOPPOPPP PEASGSWGGPPPPPPGNNATSNSWSSGPOPA TPKDEEPSGWEEPSPGSISKKMDIDDGTSAWG DPNSYNYXNNIVLWDKNSQGGPAPREEPLLPTP MTSKSASDSKSNOGOGWGESDGPVTGARIPS WEEEEDGGVWNTTGSGGSASSHNSASWGGG GKKGMKCSLKGGNNDSWMNPLAKQFSNSGG GKKGMKCSLKGGNNDSWMNPLAKQFSNSGG GKKGMKCSLKGGNNDSWMNPLAKQFSNSGG GKKGMKCSLKGGNNDSWMNPLAKQFSNSGG GKKGMKCSLKGGNNDSWMNPLAKQFSNSGG GKKGMKCSLKGGNNDSWMNPLAKQFSNSGG GKKGMKCSLKGGNNDSWMNPLAKQFSNSGG GKKGMKCSLKGGNNDSWMNPLAKQFSNSGG GKKGMKCSLKGGNNDSWMNPLAKQFSNSGG GKKGMKCSLKGGNNDSWMNPLAKQFSNSGG GKKGMKCSLKGGNNDSWMNPLAKQFSNSGG GKKGMKCSLKGGNNDSWMNPLAKQFSNSGG GKKGMKCSLKGGNNDSWMNPLAKQFSNSGLS GRADAAGALVGGPGCQCQC QCQQQQQQQQQQCGGMKHSFSFSWYGF PHDNMVPPALAVGGTCNGGSPY NQFDIJFDGTLGGHTOFAGDSWLAKASPPTINK IGKSSSASWPFFPGPQGVPWKGIQNIDPESIP YVTPGSVLGGTATSFIVOTDEHQLLRDNTTGS NSSCLAWAGGTALHMCVLGNTTIL AEF ATDDEVSRLAAAGTALHMCVLGNTTIL AEF ATDDEVSRLAAAGTALHMCVLGNTTIL AEF ATDDEVSRLAAAGTALHMCVLGNTTIL AEF ATDDEVSRLAAAGTALHMCVLGNTTIL AEF ATDDEVSRLAAAGTALHMCVLGNTTIL AEF ATDDEVSRLAAAGTALHMCVLGNTTIL AEF ATDDEVSRLAAAGTALHMCVLGNTTIL AEF ATDDEVSRLAAAGTALHMCVLGNTTIL AEF ATDDEVSRLAAAGTALHGCVLGNTSSLWSSAGGSSADLAAGALWGPPTYSSLWGYPTVED PHRMGSPAPLLFGGLIGGGSDSI VPWRRQDGGCLSQUGVTTQALTVELARWTILLLSFTF LPPSSLPPFLQUVDSSSSACTLDSFPFPLAARWDS QCGFKDBGPQTLTQALTVELARWTILLLS TAMYGAGAPPLLALCHVDGRVPFRPSSALLT ELTKLLLCASFSLLVGWGG			ļ				DMEEWPCIASKDTESSSENTTDNNSASNPGSE
GODRIGESGESTNYPNNNSAWPALVQEGTS RKGALETDNSNSSAQVETQSTSREQQSKME NAGVPFVVSGREQAGHNTDGPKRGNTNSL NISSPNPTENKGMPFGHGNTSSTAPBQ STGDRKTGSVGSWGAARGPSGTDTVSGQSNS GNNGNNGKREDSWG AVGKTGSKNDS WDNNNRSTGGSWNFGPQDSNDNKWGEGNK MTSGVSQGEWKQPTGSBLGGEWSGPNOPN STGAWDNQKGHPLLENQGNAQAPCWGRSS STOSPLGGSTGSNHKAGEGNK MTGGVSQGEWKQPTGSSSPLHNSGRRSS RPTHIPDCQAVLQTLLSRTDLDPRVLSNTGWG QTQIKQDTVWDIFEVPRPEGKSDKGTEGWSS AATQTKNSGGWGDAPSQSNQMKSGWGELS ASTEWKDPKNTGGWNDYRNNNSSNWGGGR PDEKTTSSWNEPNSKLOGWGGGROPNOGWS SGIKNGWGEEVPOTTGNSNWESGASRYVSGWG GEGQNEIGTWGNGNASLASKGGWEDCKRS. PAWNETGRQPNSWNKQHQQQPQPPPQPPPPQ PEASGSWGGPPPPFGGNYRSNNSWSSASRYVSGWG DPNSYNYKNYALWDENSQGGPAPREPNLPTP MTSKSASDSKSMQDGWGESDOPVYGARHPS WEEEDGGVWNTTGSGSGASSHNSASWGQG GKKQMKCSLKGGNNDSWMNPLAKAGNSNMG LLSQTEDNFSSKMDLSVGSLSDKKFDVDKAR MNLGPFNDIMRKDRSGFRPNSKDLMTTDS GPYPEKGGSVONTGSGSGASSHNSASWGQG GKKQMKCSLKGGNNDSWMNPLAKQPSNMG LLSQTEDNFSSKMDLSVGSLSDKKFDVDKAR MNLGPFNDIMRKDRSGFRPNSKDLMTTDS GPYPEKGGSFGLGFNSTASGLHTTVOQLN SSPSLRAQVPPOFISQVSASNLKGPDVDKAR MNLGPFNDIMRKDRSGFRPNSKDLMTTDS GPYPEKGGSFGLGFNSTASGLHTVOQLN SSPSLRAQVPPOFISQVSASNLKGPQBQCA RMVSALQQQQQOLQNGRKGAPPSGCHGFNSGLSP PHLDNMVPNALNVGLPDLQTKGPPFGCGGF PHLDNMVPNALNVGLPDLQTKGPFPGTGGGF PHLDNMVPNALNVGLPDLQTKGGPFPYGKGF PHLDNMVPNALNVGLPDLQTKGPFPGTGGF SGGMDYGWGKKGATSPRYVVFK MSSLNTSLPSFGAWPYSASDNLFNVHSTSAK PPDYKSTWSPPGFQTATHLNCTLGGTTLLAFT LIRYSTKQEAARAGATALTMCTLLAFT LIRYSTKQEAARAGATALTMCTLLAFTILLAFT LIRYSTKQEAARAGATALTMCTLLAFTILLAFT LIRYSTKQEAARAGATALTMCTLLAFTILLAFT LIRYSTKQEAARAGATALTMCTLLAFTILLAFT ALTDDEVSRFLAQAQPPTFPAATPSAPAAGWGS LETGQNQSDPCGARLAFGGSTSPVANVALLL LTPODGSTLRTICMQHGPLLTPHLNLTQTTA LIRYSTKQEAARAGATALTMCTLLAFTILLAFT ALTDDEVSRFLAQAQPPTFPAATPSAPAAGWGS LETGQNQSDPCGARLAFGGSTSPVANVALLL LTPODGSTLRTICMQHGPLLTPHLNLTTQTTA LIRYSTKQEAARAGATALTMCTLLAFTILLAFT ALTDDEVSRFLAQAQPTFTPAATPSAPAAGWGS BATTPLPRPFPGLAAGAPTPTAATPSAPAAGWGS LETGQNQSDPCAGARLAFGGSTGPAATPSAPAGT PHRMGSPAPLLFGGGTSLGVETTLTLLAFT LIRYSTKQEAARAGATALTMCTLAFTTLLAFT LIRYSTKQEAARAGATALTMCTLAFTTLLAFT LIRYSTKQEAARAGATALTMCTLAFTTLLAFT LIRYSTKQEAARAGATALTMCTLAFTTLLAFT LIRYSTKQEAARAGATALTMCTLAFTTLAFT	1	l	İ			}	KSTLPGSTTSNKGKGSQCQSASSGNECNLGV
RKGALETDNSNSSAQVSTVGGTSREQQŠKME NAGVNFVVSGREQAQHTOPGFKNGTNISL NLSSPNFMENKGMPFGMGLGNTSRSTDAPSQ STGDRKTGSVGSWGAAGAGSVQKSTGSKNDS WDNNNSTGGSWNFPGMGLGNTSRSTDAPSQ STGDRKTGGSWNFPGDSDDNKWGEGNK MTSGVSQGEWKQPTGSDELKIGEWSGPNQFN STGAWDNQKGHPLLENQGNAQAFWGRSS SSTGSEVEGSTGSNHAAGASSDKSTGSKNDS WDNNNSTGGSWNFPGDSDDNKWGEGNK MTSGVSQGEWKQPTGSDELKIGEWSGPNQFN STGAWDNQKGHPLLENQGNAQAFWGRSS SSTGSEVEGSTGSNHAAGASSDSINNSGRSY RPHPDCQAVLQTLLSRTDLDPRVLSNTGWG QTQIKQDTVVDIEEVPFFEGKSDKGTEGWES AATQTKNSGGWGDAPSQNGMKSGWGELS ASTEWKDPFFGKSDKGTEGWES AATQTKNSGGWGDAPSQNGMKSGWGELS ASTEWKDPFFGKSDKGTEGWES AATQTKNSGGWGDAPSQNGMKSGWGELS ASTEWKDPFSKDQGWGGGROPNQGWS SGKNGWGEEVDOTKNSNWESSASKPVSGWG EGGNEIGTWGNGGNASLASKGGWDEKSR PAWNETGREPNSWNENDESSASKPVSGWG EGGNEIGTWGNGGNASLASKGGWDEKSR PAWNETGREPNSWNENDESSASKPVSGWG EGGNEIGTWGNGGNASLASKGGWDEKSR PAWNETGREPNSWNENDESSASKPVSGWG EGGNEIGTWGNGGNASLASKGGWDEKSR PAWNETGREPNSWNENDESSASKPVSGWG EGGNEIGTWGNGGASASHNASSWGQG GKKOMKCSKGSNDSWMPILAKGPSNMG LLSQTEDNPSSKMDLSVGSGSASHNASSWGGG GKKOMKCSLKGGNNDSWMPILAKGPSNMG LLSQTEDNPSSKMDLSVGSLSDKKFDDVKTAAHP WEEEDDGGVWNTTGSGGSASSHNASSWGGG GKKOMKCSLKGGNNDSWMPILAKGPSNMG LLSQTEDNPSSKMDLSVGSLSDKKFDDVKTAAHP WEEEDDGGVWNTTGSGASASHNASWGGG GKKOMKCSLKGGNNDSWMPILAKGPNSGLPT WEEGDGGVWNTTGSGASSANGASGGPPPPKCDLACQL LLQQQQQQLLQNQRKISQAXGCLPDFPGYGAGG GKYOMKCSLKGGNNDSWMPILAKGPNSGLPT WEEGDGGVWNTGCKAGTESRFKQWTSMME GLPSVATQEANHIKNGAIVAPGKTRGGSPY NGFDIPGDTLGGTTGFAGSNLPASKPTNK IGSKSSNASWPPEPOPGVPWKGGNDFESDP YVTTGSVLGGTATSPTUDTHQLLRDNTTGS NSSLNTSLPSPGAWPYSASDNSFTNVHSTSAK FPDYKSTWSPPDIGHNTPILAKFASWYNTHSTSAK FPDYKSTWSPPDIGHNTPILAKFASWYNTHSTSAK FPDYKSTWSPPDIGHNTPILAKFASWYNTHSTAK FPDYKSTWSPPDIGHNTPILAKFASWYNTHSTAK FPDYKSTWSPPDIGHNTPILAKFASWYNTHSTAK FPDYKSTWSPPDIGHNTPINKYNSSNIPTNVHSTSAK FPDYKSTWSPPDIGHNTPINKYNSSNIPTNVHSTSAK FPDYKSTWSPPDIGHNTPINKYNSSMPTNVHSTSAK FPDYKSTWSPPDIGHNTPINAKFASWYNTHSTAK FPDYKSTWSPPDIGHNTPINAKFASWYNTHSTAK FPDYKSTWSPPDIGHNTPINAKFASWYNTHSTAK FPDYKSTWSPPDIGHNTPINAKFASWYNTHALLE FLOWGOSGADLAGASLWGPPYYSSSLWGVPTVED PHRMGSPAPLJOGDLLGGGSDSI LTTLLLCHAGGSTGLGGWSSA GGSGADLAGASLWGPPYYSSSLWSSACTLDSFTPILAPWD PQCGF							WKSDPKAKSVQSSNSTTENNNGLGNWRNVS
NAGYNFVYSGREQAÖIHNTDÖRENGNTRSSTDAPSQ STÖDRKTGSVGSWGAARGPSGTDYSGOSNS GNNGNNGKREDSWG AS SYCKSTGSKNDS WDNNNRSTGGSWNFGPQDSNDNKWGEGNK MTSGVSQGWKQPTGSDELKIGEWSGPNQPN SSTGAWDNQKGHPLLENQMAQAPCWGRSS STGSVEGQSTGSNHKAGSSDSINSGRRSY RPTHPDCQAVIQTLLSKTDLDPRVLSNTGWG QTQIKQDTVWDIBEVPRFEGKSDKGTEGWES AATOTKNSGGWGDAPSQMKSGWGELS ASTEWKDPKNTGGWDYYKNNNSSNWGGGR PDEKTPSSWNEPSRLOQ WGGGRQPNQGWS GGKNGWGEEVDDTKNSNWESSASKPVSGWG EGGONEIGTWGNGNAS KAGGWEDCKRS. PAWNETGRQPNSWNKQHQQQQPQPPPP PEAAGSWGGPPPPPGNTSSSWSGGPQPA TPKDEPSGWERPSQGSTRSKMDDGTSAWG DPNSYNYKNVALWDENSSWSSGPQPA TPKDEPSGWERPSQGSASSHNASWQGG GKKQMKCSLKGGNNDSWMPLAKQFSNMG LLSQTEDNPSSKMDLSVGSLSDKKFDDTSAWG MANLGDFNDINKDRSGFRPPNSKDMGTTDS GYPTEKGGSIGLFGNSTAQSRGLHTPVQFLN SSPSLRAQVPPQFSPQVSASMLKQPPNSGLSP GLTNWGPQLSPQQIAMLSQLFQPTOFLACQL LLQQQQQQOQQRQFGMKHSPSPPVGSGF SGGMDYGMVAGKEAGTESRKQWTSMMG GLPSVATQGALLQQQQQQOQQRQFGMKHSPSPPVGKF PHLDNMVPALNVGLPJCHACKGSPP NQFDIPGDTLGGHTGPAGDSWLPAKSPPTNK IGSKSNASWPPERQGVFWKGRJONEGSPP NQFDIPGDTLGGHTGPAGDSWLPAKSPPTNK IGSKSNASWPPERQGVFWKGRJONEGSPP NQFDIPGDTLGGHTGPAGDSWLPAKSPPTNK IGSKSNASWPPERQGVFWKGRJONEGSPP NQFDIPGDTLGGHTGPAGDSWLPAKSPPTNK IGSKSNASWPPERQGVFWKGRJONEGSPP NQFDIPGDTLGGHTGPAGDSWLPAKSPPTNK IGSKSNASWPPERQGVFWKGRJONEGSPP NQFDIPGDTLGGHTGPAGDSWLPAKSPPTNK IGSKSNASWPPERQGVFWKGRJONEGSPP NQFDIPGDTLGGHTGPAGDSWLPAKSPPTNK IGSKSNASWPPERQGSVRPSYWLVLHN LTTQIDGSTLRTICMQHGPLLTTHLNTLTQGTA LIRYSTKQEAKAQTALTH-LSGSTGLGQWSSSA GGSSGADLAGASLWGPPPNSSSLWGVPTVED PHRMGSPALLFGDLLGGGSDSI LETGYNGARAAGTUALTH-LNTQGTA LIRYSTKGEAKAQTALTH-LAGGSTSL LIRYSTKGEAKAQTALTH-LAGGSTSL LIRYSTKGEAKAQTALTH-LAGGSTSL LIRYSTKGEAKAQTALTH-LAGGSTSL LIRYSTKGEAKAQTALTH-LNTLTQGTA LIRYSTKGEAKAQTALTH-LNTGGTSTGLGQWSSSA GGSSGADLAGASLWGPPPNSSSLWGVPTVED PHRMGSPAPLLFFOLLIGGGSDSI PRYMGSPAPLLFFOLLIGGGSDSI PRYMGSPAPLLFFOLLIGGGSDSI PRYMGSPAPLLFFOLLIGGGSDSI PRYMGSPAPLLFFOLLIGGGSDSI PRYMGSPAPLLFFOLLIGGGSDSI PRYMGSPAPLLFFOLLIGGGSDSI PRYMGSPAPLLFFOLLIGGGSDSI PRYMGSPAPLLFFOLLIGGGSDSI PRYMGSPAPLLFFOLLIGGSDSI PRYMGSPAPLLFFOLLIGGSDSI PRYMGSPAPLLFFOLLIGGSDSI PRYMGSPARAGWQAPPPWRQAA							GQDRIGPGSGFSNFNPNSNPSAWPALVQEGTS
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YVTPGSVLGGTATSPIVDTDHQLLRDNTTGS NSSLNTSLPSPGAWPYSASDNSFTNVHSTSAK FPDYKSTWSPDPIGHNPTHLSNKMWKNHISS RNTTPLPRPPPGLTNPKPSSPWSSTAPRSVRG WGTQDSRLASASTWSDGGSVRPSYWLVLHN LTPQIDGSTLRTICMQHGPLLTFHLNLTQGTA LIRYSTKQEAAKAQTALHMCVLGNTTILAEF ATDDEVSRFLAQAQPPTPAATPSAPAAGWQS LETGQNQSDPVGPALNLFGGSTGLGQWSSSA GGSSGADLAGASLWGPPNYSSSLWGVPTVED PHRMGSPAPLLPGDLLGGGSDSI  1049 2399 A 8748 200 1387 VPWKRQDEQLSLQVETLYLDSPAVIHLLSPTF LPPSSLPPFLQIVDSSSSACTLDSFFPFLAPWDS PQDCGFKDHQPLTLQALTVELARWTLMLLLS TAMYGAHAPLLALCHVDGRVPFRPSSAVLLT ELTKLLLCAFSLLVGWQAWPQGPPPWRQAA		1	1				
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FPDYKSTWSPDPIGHNPTHLSNKMWKNHISS RNTTPLPRPPPGLTNPKPSSPWSSTAPRSVRG WGTQDSRLASASTWSDGGSVRPSYWLVLHN LTPQIDGSTLRTICMQHGPLLTFHLNLTQGTA LIRYSTKQEAAKAQTALHMCVLGNTTILAEF ATDDEVSRFLAQAQPPTPAATPSAPAAGWQS LETGQNQSDPVGPALNLFGGSTGLGQWSSSA GGSSGADLAGASLWGPPNYSSSLWGVPTVED PHRMGSPAPLLPGDLLGGGSDSI  1049 2399 A 8748 200 1387 VPWKRQDEQLSLQVETLYLDSPAVIHLLSPTF LPPSSLPFFLQIVDSSSSACTLDSFFPFLAPWDS PQDCGFKDHQPLTLQALTVELARWTLMLLLS TAMYGAHAPLLALCHVDGRVPFRPSSAVLLT ELTKLLLCAFSLLVGWQAWPQGPPPWRQAA	1	1	ł				
RNTTPLPRPPPGLTNPKPSSPWSSTAPRSVRG WGTQDSRLASASTWSDGGSVRPSYWLVLHN LTPQIDGSTLRTICMQHGPLLTFHLNLTQGTA LIRYSTKQEAAKAQTALHMCVLGNTTILAEF ATDDEVSRFLAQAQPPTPAATPSAPAAGWQS LETGQNQSDPVGPALNLFGGSTGLGQWSSSA GGSSGADLAGASLWGPPNYSSSLWGVPTVED PHRMGSPAPLLPGDLLGGGSDSI  1049 2399 A 8748 200 1387 VPWKRQDEQLSLQVETLYLDSPAVIHLLSPTF LPPSSLPPFLQIVDSSSSACTLDSFFPFLAPWDS PQDCGFKDHQPLTLQALTVELARWTLMLLLS TAMYGAHAPLLALCHVDGRVPFRPSSAVLLT ELTKLLLCAFSLLVGWQAWPQGPPPWRQAA		I	1				
WGTQDSRLASASTWSDGGSVRPSYWLVLHN LTPQIDGSTLRTICMQHGPLLTFHLNLTQGTA LIRYSTKQEAAKAQTALHMCVLGNTTILAEF ATDDEVSRFLAQAQPPTPAATPSAPAAGWQS LETGQNQSDPVGPALNLFGGSTGLGQWSSSA GGSSGADLAGASLWGPPNYSSSLWGVPTVED PHRMGSPAPLLPGDLLGGGSDSI 1049 2399 A 8748 200 1387 VPWKRQDEQLSLQVETLYLDSPAVIHLLSPTF LPPSSLPPFLQIVDSSSSACTLDSFFPFLAPWDS PQDCGFKDHQPLTLQALTVELARWTLMLLLS TAMYGAHAPLLALCHVDGRVPFRPSSAVLLT ELTKLLLCAFSLLVGWQAWPQGPPPWRQAA							
LTPQIDGSTLRTICMQHGPLLTFHLNLTQGTA LIRYSTKQEAAKAQTALHMCVLGNTTILAEF ATDDEVSRFLAQAQPPTPAATPSAPAAGWQS LETGQNQSDPVGPALNLFGGSTGLGQWSSSA GGSSGADLAGASLWGPPNYSSSLWGVPTVED PHRMGSPAPLLPGDLLGGGSDSI  1049 2399 A 8748 200 1387 VPWKRQDEQLSLQVETLYLDSPAVIHLLSPTF LPPSSLPPFLQIVDSSSSACTLDSFFPFLAPWDS PQDCGFKDHQPLTLQALTVELARWTLMLLLS TAMYGAHAPLLALCHVDGRVPFRPSSAVLLT ELTKLLLCAFSLLVGWQAWPQGPPPWRQAA							
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ATDDEVSRFLAQAQPPTPAATPSAPAAGWQS LETGQNQSDPVGPALNLFGGSTGLGQWSSSA GGSSGADLAGASLWGPPNYSSSLWGVPTVED PHRMGSPAPLLPGDLLGGGSDSI  1049 2399 A 8748 200 1387 VPWKRQDEQLSLQVETLYLDSPAVIHLLSPTF LPPSSLPPFLQIVDSSSSACTLDSFFPFLAPWDS PQDCGFKDHQPLTLQALTVELARWTLMLLLS TAMYGAHAPLLALCHVDGRVPFRPSSAVLLT ELTKLLLCAFSLLVGWQAWPQGPPPWRQAA		1	1				
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PQDCGFKDHQPLTLQALTVELARWTLMLLLS TAMYGAHAPLLALCHVDGRVPFRPSSAVLLT ELTKLLLCAFSLLVGWQAWPQGPPPWRQAA	1049	2333	^	0/40	200	1501	
TAMYGAHAPLLALCHVDGRVPFRPSSAVLLT ELTKLLLCAFSLLVGWQAWPQGPPPWRQAA	1	I					
ELTKLLLCAFSLLVGWQAWPQGPPPWRQAA		1	[				
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PFALSALLYGANNNLVIYLORYMDPSTYOVI.		1	1			:	
	L ·	}	<u></u>	J i			PFALSALLYGANNNLVIYLQRYMDPSTYQVL

SEQ ID NO: of nucl- cotide seq- uence	SEQ ID NO: of peptide scq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  SNLKIGSTAVLYCLCLRHRLSVRQGLALLLL MAAGACYAAGGLQVPGNTLPSPPPAAAASP MPLHITPLGLLLILYCLISGLSSVYTELLMKR QRLPLALQNLFLYTFGVLLNLGLHAGGGSGP GLLEGFSGWAALVVLSQALNGLLMSAVMKH GSSITRLFVVSCSLVVNAVLSAVLLRLQLTAA FFLATLLIGLAMRLYYGSR
1050	2400	A	8758	3	1660	WVSSMGFEELLEQVGGFGPFQLRNVALLALP RVLLPLHFLLPIFLAAVPAHRCALPGAPANFS HQDVWLEAHLPREPDGTLSSCLRFAYPQALP NTTLGEERQSRGELEDEPATVPCSQGWEYDH SEFSSTIATESQWDLVCEQKGLNRAASTFFFA GVLVGAVAFGYLSDRFGRRRLLLVAYVSTLV LGLASAASVSYVMFAITRTLTGSALAGFTIIV MPLELEWLDVEHRTVAGVLSSTFWTGGVML LALVGYLIRDWRWLLLAVTLPCAPGILSLWW VPESARWLLTQGHVKEAHRYLLHCARLNGR PVCEDSFSQEAVSKVAAGERVVRRPSYLDLF RTPRLRHISLCCVVVWFGVNFSYYGLSLDVS GLGLNVYQTQLLFGAVELPSKLLVYLSVRYA GRRLTQAGTLLGTALAFGTRLLVSSDMKSWS TVLAVMGKAFSEAAFTTAYLFTSELYPTVLR QTGMGLTALVGRLGGSLAPLAALLDGVWLS LPKLTYGGIALLAAGTALLPETRQAQLPETI QDVERKSAPTSLQEEEMPMKQVQN
1051	2401	A	8759	515	1625	EIRTPVAVSSAPSGDSEGDEEETTQDEVSSHTS EEDGGVVKVEKELENTEQPVGGNEVVEHEV TGNLNSDPLLELCQCPLCQLDCGSREQLIAHV YQHTAAVVSAKSYMCPVCGRALSSPGSLGR HLLIHSEDQRSNCAVCGARFTSHATFNSEKLP EVLNMESLPTVHNEGPSSAEGKDIAFSPPVYP AGILLVCNNCAAYRKLLEAQTPSVRKWALRR QNEPLEVRLQRLERERTAKKSRRDNETPEERE VRRMRDREAKRLQRMQETDEQRARRLQRDR EAMRLKRANETPEKRQARLIREREAKRLKRR LEKMDMMLRAQFGQDPSAMAALAAEMNFF QLPVSGVELDSQLLGKMAFEEQNSSSLH
1052	2402	A	8763	1106	70	RHGHGGRDRRGGGRVARPGGLGRYPGRGAA ASLVFVPTRRRSGPSGTASVAAMAYHSGYGA HGSKHRARAAPDPPPLFDDTSGGYSSQPGGY PATGADVAFSVNHLLGDPMANVAMAYGSSI ASHGKDMVHKELHRFVSVSKLKYFFAVDTA YVAKKLGLLVFPYTHQNWEVQYSRDAPLPP RQDLNAPDLYIPTMAFITYVLLAGMALGIQK RFSPEVLGLCASTALVWVVMEVLALLLGLYL ATVRSDLSTFHLLAYSGYKYVGMILSVLTGL LFGSDGYYVALAWTSSALMYFIVRSLRTAAL GPDSMGGPVPRQRLQLYLTLGAAAFQPLIIY WLTFHLVR
1053	2403	A	8768	2	712	RPPRVWYPELRELSAAAPRWSHRTAPGIMVF YFTSSSVNSSAYTIYMGKDKYENEDLIKHGW PEDIWFHVDKLSSAHVYLRLHKGENIEDIPKE VLMDCAHLVKANSIQGCKMNNVNVVYTPW SNLKKTADMDVGQIGFHRQKDVKIVTVEKK VNEILNRLEKTKVERFPDLAAEKECRDREER NEKKAQIQEMKKREKEEMKKKREMDELRSY SSLMKVENMSSNQDGNDSDEFM
1054	2404	Α	8769	344	527	REATTLACRNSCWVFSRCSLGACKPTVCSMP SLSRQGSQTLCLRLAEYCMESVDSQRLLLS

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
1055	2405	A	8770	430	1104	QQESPAAGAARMNCKEGTDSSCGCRGNDEK KMLKCVVVGDGAVGKTCLLMSYANDAFPEE YVPTVFDHYAVTVTVGGKQHLLGLYDTAGQ EDYNQLRPLSYPNTDVFLICFSVVNPASYHNV QEEWVPELKDCMPHVPYVLIGTQIDLRDDPK TLARLLYMKEKPLTYEHGVKLAKAIGAQCYL ECSALTQKGLKAVFDEAILTIFHPKKKKKRCS EGHSCCSII
1056	2406	A	8773	261	332	NPRIQLSGNSCCAGSCRVWLSEQ
1057	2407	A	8778	3	477	PAGIRHEQARGADRMGKCRGLRTARKLRSH RRDQKWHDKQYKKAHLGTALKANPFGGAS HAKGIVLEKVGVEAKQPNSAIRKCVRVQLIK NGKKITAFVPNDGCLNFIEENDEVLVAGFGR KGHAVGDIPGVRFKVVKVANVSLLALYKGK KERPRS
1058	2408	A	8808	171	881	PGLSQEPSGSMETVVIVAIGVLATIFLASFAAL VLVCRQRYCRPRDLLQRYDSKPIVDLIGAME TQSEPSELELDDVVITNPHIEAILENEDWIEDA SGLMSHCIAILKICHTLTEKLVAMTMGSGAK MKTSASVSDIIVVAKRISPRVDDVVKSMYPPL DPKLLDARTTALLLSVSHLVLVTRNACHLTG GLDWIDQSLSAAEEHLEVLREAALASEPDKG LPGPEGFLQEQSAI
1059	2409	Ā	8809	246	757	MRLQGAIFVLLPHLGPILVWLFTRDHMSGWC EGPRMLSWCPFYKVLLLVQTAIYSVVGYASY LVWKDLGGGLGWPLALPLGLYAVQLTISWT VLVLFFTVHNPGLALLHLLLLYGLVVSTALI WHPINKLAALLLLPYLAWLTVTSALTYHLWR DSLCPVHQPQPTEKSD
1060	2410	A	8810	304	381	PKLSVYPLQSHHCLSEPFQSLVCCLA
1061	2411	A	8820	1673	848	SCKTENLLEMWWFQQGLSFLPSALVIWTSAA FIFSYITAVTLHHIDPALPYISDTGTVAPEKCLF GAMLNIAAVLCIATIYVRYKQVHALSPEENVI IKLNKAGLVLGILSCLGLSIVANFQKTTLFAA HVSGAVLTFGMGSLYMFVQTILSYQMQPKIH GKQVFWIRLLLVIWCGVSALSMLTCSSVLHS GNFGTDLEQKLHWNPEDKGYVLHMITTAAE WSMSFSFFGFFLTYIRDFQKISLRVEANLHGL TLYDTAPCPINNERTRLLSRDI
1062	2412	A	8824	1	763	GGAPPASVPARESPVSGAQGSSRTRGHKRAA GARAPQLCSSWQRRSAPAMSRGLQLLLLSCA YSLAPATPEVKVACSEDVDLPCTAPWDPQVP YTVSWVKLLEGGEERMETPQEDHLRGQHYH QKGQNGSFDAPNERPYSLKIRNTTSCNSGTYR CTLQDPDGQRNLSGKVILRVTGCPAQRKEET FKKYRAEIVLLLALVIFYLTLIIFTCKFARLQSI FPDFSKAGMERAFLPVTSPNKHLGLVTPHKT ELV
1063	2413	A.	8826	147	627	CETSTSSAGHAPCRHAAQGPPAEPTGLRLCSE HQRLHAWPPGPRRPSLWPPKNGKWHSGKRT AGGRPQRRPSRRQSQRPSAWSGSPRMHSPGQ KCSLMCPHRSQDSLSTAIFQRSPGANTGRALH CVLSKEMKSVQRSLGLSRIHLQSKRKIIHFVL TR
1064	2414	A	8835	2982	1869	LKDTLKSQMTQEASDEAEDMKEAMNRMIDE LNKQVSELSQLYKEAQAELEDYRKRKSLEDV TAEYIHKAEHEKLMQLTNVSRAKAEDALSE MKSQYSKVLNELTQLKQLVDAQKENSVSITE HLQVITTLRTAAKEMEEKISNLKEHLASKEVE

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Scrine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  VAKLEKQLLEEKAAMTDAMVPRSSYEKLQS SLESEVSVLASKLKESVKEKEKVHSEVVQIRS EVSQVKREKENIQTLLKSKEQEVNELLQKFQ QAQEELAEMKRYSESSSKLEEDKDKKINEMS KEVTKLKEALNSLSQLSYSTSSSKRQSQQLEA LQQVKQLQNQLAECKKQHQEVISVYRMHL LYAVQGQMDEDVQKVLKQILTMCKNQSQK K
1065	2415	A	8841	3	663	AAATAASLSPRGCRLRTPSSDVGPSRAPPPSA APLPTGRAQMSPSGRLCLLTIVGLILPTRGQTL KDTTSSSSADATIMDIQVPTRAPDAVYTELQP TSPTPTWPADETPQPQTQTQQLEGTDGPLVT DPETHKSTKAAHPTDDTTTLSERPSPSTDVQT DPQTLKPSGFHEDDPFFYDEHTLRKRGLLVA AVLFITGIIILTSGKCRQLSRLCRNHCR
1066	2416	A	8853	3806	2204	FVGEQEGGCEAGAGRGAQTYPGEAGERWFG RRRRRGRVVSRKKMSLKSERRGIHVDQSDLL CKKGCGYYGNPAWQGFCSKCWREEYHKAR QKQIQEDWELAERLQREEEEAFASSQSSQGA QSLTFSKFEEKKTNEKTRKVTTVKKFFSASSR VGSKKEIQEAKAPSPSINRQTSIETDRVSKEFIE FLKTFHKTGQEIYKQTKLFLEGMHYKRDLSIE EQSECAQDFYHNVAERMQTRGKVPPERVEKI MDQIEKYIMTRLYKYVFCPETTDDEKKDLAI QKRIRALRWVTPQMLCVPVNEDIPEVSDMVV KAITDIIEMDSKRVPRDKLACITKCSKHIFNAI KITKNEPASADDFLPTLIYIVLKGNPPRLQSNI QYITRFCNPSRLMTGEDGYYFTNLCCAVAFIE KLDAQSLNLSQEDFDRYMSGQTSPRKQEAES WSPDACLGVKQMYKNLDLLSQLNERQERIM NEAKKLEKDLIDWTDGIAREVQDIVEKYPLEI KPPNQPLAAIDSENVENDKLPPPLQPQVYAG
1067	2417	A	8855	1372	1513	SNMREVGCGWLVPVIPAFWEAEVGGSLEARS LRQAWATKQDPISKKK
1068	2418	A	8856	<b>15</b> 30	1583	PCRPGMECNSMISVHCNL
1069	2419 2420	A	8857	1530 293	1583 1675	PCRPGMECNSMISVHCNL  PYPQGGYPQGPYPQEGYPQGPYPQGGYPQGP YPQSPFPPNPYGQPQVFPGQDPDSPQHGNYQ EEGPPSYYDNQDFPATNWDDKSIRQAFIRKVF LVLTLQLSVTLSTVSVFTFVAEVKGFVRENV WTYYVSYAVFFISLIVLSCCGDFRKHPWNL VALSVLTASLSYMVGMIASFYNTEAVIMAVG ITTAVCFTVVIFSMQTRYDFTSCMGVLLVSM VVLFIFAILCIFIRNRILEIVYASLGALLFTCFLA VDTQLLLGNKQLSLSPEEYVFAALNLYTDIINI FLYILTIIGRAKE*PSSSSLCPLRWHGWPGPCP WHGSASCTSPLSCPQAQPREKDASLQPSCMY TADTSIWTRCGHSMAPLVLPPPPRGTKATFPC HLLSTHCCMSPVCQPTPGTGGSTRSRGEGLSQ EVRVHVFPPVPAPQPGVEHPSPPPHPPGVLPS GDMRSGGLIPVLSPE
1071	2421	A	8868	2	358	ARGNTLYHLPRLCRKLNLRWFSASTLYDVQH DDKMGSNTFFKRNDCRYVMISCKADMAYDN VRHPFMI*SIKLIMEETYLNIIKAVYDRPTASII LNGEKLKVFPVRSGT*QGCSVWP
1072	2422	A	8870	33	658	MESVLSKYEDQITIFTDYLEEYPDTDELVWIL GKQHLLKTEKSKLLSDISARLWFTYRRKFSPI GGTGPSSDAGWGCMLRCGQMMLAQALICRH LGRDWSWEKQKEQPKEYQRILQCFLDRKDC

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						CYSIHQMAQMGVGEGKSIGEWVLGPNTV\AQ GV*KNLA\LFDEW\NSLGLVYVSM\DNPSGSIA RFPKKLCRVLPL\SADTAGLTGP
1073	2423	A	8879	146	412	DFSV*GDVDIEVTCPICLQLLTEPLSLNCGLRL *QVCITA*IKESVIISGG*SSSPVCHTTFQPANL RTSRYLPT*SIKSLGPDEPQEG
1074	2424	A	8884	67	435	HLQGRSIRTLQLTGENEKNCEVSERIRRSGPW KEISFGDYICHTFQGDCWADRSPLHEAAAHG RLLALKTLIAQGVNVNLWTL/DRVSSLHEACL *GPVACAKPYWKMVPRHGGTVTGPPLLMV
1075	2425	A	8896	1294	248	RSGDRNGLTHQLGGLSQGSRNQSYRSRSRSR SRERPSAPRGIPFASASSSVYYGSYSRPYGSDK PWPSLLDKEREESLRQKRLSERERIGELGAPE VWGLSPKNPEPDSDEHTPVEDEEPKKSTTSAS TSEEEKKKKSSRSKERSKKRRKKKSSKRKHK KYSEDSDSDSDSETDSSDEDNKRRAKKAKKK EKKKKHRSKKYKKKRSKKSRKESSDSSSKES QEEFLENPWKDRTKAEFSDLIGPEAPKTLTS QDDKPLNYGHALLPGEGAAMAEYVKAGKRI PRRGEIGLTR*RNCHHLNAQVM**VVSRHRR MEAVRTAKREPESTVLMRREPLHPFNPRRET KERE
1076	2426	A	8899	146	789	GRSTEAEKEPAFDERTGKGRRLPRAGEFHG*E *APGPGPRSFQVSRKMPEE\PPGARKHPFSGKS FYLDLPAGKNLQFLTGAIQQLGGVIEGFLSKE VSYIVSSRREVKAESSGKSHRGCPSPSPSEVR VETSAMVDPKGSHPRPSRKPVDSVPLSRGKE LLQKAIRNQK**CTVQQLSHCRLY\GEKTTAK RSQREHVQQQSQEHGKWPDLKGPR
1077	2427	A	8901	352	3	AKIGAYKYIQELWRKKQSDVMHFLLRVRCW QYPALHRAGTEWQLSALHRAPRSTQPDKAC RLGYKAKQGYIIYRICVRRGGWKCPVPKAVT \YGKPVHHGVN*LKFAQSLQSVAEEQ
1078	2428	A	8905	536	781	ACPAENREVPEMAAGQAPHAGPGAGPGQPA PALPFAATPGSRGQALCRGGRRRQHLHGPLH RP*QAAPALHAGCQLAPHPPT
1079	2429	A	8912	121	376 -	NLIWKLCVTERRLVILDNYDLASE/YEANKYI CNRIIQFKPGQDKYFTLGLPTGSTPL*CYPKLI EYNKNGHLSFKYVKTFSMDEY
1080	2430	A	8920	381	1788	SSESPSDPGRMAMTWIVFSLWPLTVFMGHIG GHSLFSCEPITLRMCQDLPYNTTFMPNLLNHY DQQTAALAMEPFHPMVNLDCSRDFRPFLCAL YAPICMEYGRVTLPCRRLCQRAYSECSKLME MFGVPWPEDMECSRFPDCDEPYPRLVDLNLA GEPTEGAPVAVQRDYGFWCPRELKIDPDLGY SFLHVRDCSPPCPNMYFRREELSFARYFIGLIS IICLSATLFIFVTFLIDVTRFRYPERPIKCYAV WHMMVSLIFF\IGFLLEDRVACNA\SIPAQYKA STVTQGSHNKACTMLFMILYFFTMAGSVWW VILTITWFLAAVPKWGSEAIEKKALLFHASA WGIPGTLTIILLAMNKIEGDNISGVCFVGLYD VDALRYFVLAPLCLYVVVGVSLLLAGIISLNR VRIEIPL*KENQDKLVKFMIRIGVFSILYLVPLL VVIGCYFYEQAYRGIWETTWIQERC
1081	2431	A	8922	56	420	EERTKMSTGPDVKATVGDISSDGNLNVAQEE CSRKGIVDEFFPLLSN*CIWTQPQGYPQSSYG TLANFVF\CSVRHGLALILQLCNFSIYTQQMN LSIAIPAMVNNTAPPSQPNASTERPST
1082	2432	A	8923	355	1079	PFGTPSSTMAVVKNKCLMKGGKKGVKKKVV

SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-	ļ	USSN	location	corresponding to last amino	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496 914	correspondi ng to first	acid residue	M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
uence		l	914	amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
				peptide	sequence	/=possible nucleotide deletion, \=possible
	ł	ļ	İ	sequence	}	nucleotide insertion
			<del> </del>	sequence		GPFSKKDQYDVKAPAMFNIRNTGK/TLVART
						QGTQIASDGLKGLLFEVSLADLQNDEVAFRK
	ļ	ļ				FKLITEDVODKNCLTNFYGMDLTCDKICSMV
						EKWSTMIEAHVDVKTTDGYFFHLFCVGFTKK
						HNNQILKTSYA*HQQS/RQIQKKMMEIMT*EV
1	J					QTNDLKEVVNKLIPDNIGKDTEKV/CPIYPLH
						DVFIRKVKMLENPGFER\MELRGGGSSS
1083	2433	A	8948	28	385	LTWPQPHIPSCPAMSEETLQSKLAAAKKKLP
						WGAVQGSRAMSDLLLLLLDLTLLLLLMLLGF
	1					AGYSGQLAGVAVSAGSPPI/RYKFHVEPYGET
						GWLLT/ESCSISPKLCSIAVH*DNPAWF
1084	2434	A	8950	156	318	HYTPINTDTIENSENNKCW*GY*E\VGLIHHW
1005	2425	- <u>-</u> -	0056	16	412	WGGKRVQPFWKRVWQKRTLNLRV
1085	2435	A	8956	16	413	HMGQLGYFIQCWWECKRLISF\WKTI*QSPAK *TIYTSYDTAIPIS/GI/YPKRMSSKCHQETCAR
						MFILAPFTATIKGKOLTCPLVEERIDY\MWYS
						HKYYIKVKRNL*VTITH\TWVNLNILMFEIILW
						YSHKYY
1086	2436	A	8962	868	1026	H*KILQVGRAQRAHXSRL*SQLLRRLRHESHL
1000	2430	A	6702	808	1020	NPGARGCSEARLHRCTPAWTT
1087	2437	A	8985	58	330	LHVKHLGHFQLVFSEVICHCILMPVS*ELQRL
1007	2137	11	0703	30	330	*ERSVCAFHVCIQTYVCLQVYACMCVYYICM
	ł	l			}	FVYSVYGCGLCTCVCMDVYICVCVQEFL
1088	2438	A	8989	394	404	N*KWILHVNVRIQSIFF/IKRNQK/INSHELKLD
						KKFLDMMSNA*STKKHDKLD/LIKFKT/LCSA
						KYTVKRIKIHPTDLEKMLRNHLSDKD*YS/GV
	ļ	J			l	YKDLSKLNRRKTE/S*/VKKWVKDLSRYFIKE
	<u> </u>					VISMENKHKKIFSTS
1089	2439	A	8991	60	329	MALTPESPSSFPGLAATGSSVPEPPGGPNATL
	1					NSSWDSPTEPSSLEDLEATGTIGTLLSDMGVV
						GVEDNAYTLEVNSRYMRAVGIM*IHL
1090	2440	A	8996	2	351	SNITITLT*MKKYDNTFCW*GCGQIG/T/LIYC
						WQESKFIQAFWSKIQQYLA*ISIHILFDPAFLFL
1	1					GGYPGGTQSVFLTGVLVSSVFYNMKMLHTR
1001	2441		0007	07	450	LLIAALFIIVQYWKQSKDHYI
1091	2441	A	8997	97	456	YPLPVCSYLSGPRGEHWNSLGGKSSCPLPLPT LVSSRFKISKVIVVGDLSVGKTCLINR*GGAG
		1				AELGRVGPSLARWAGSRSQHLVPSQ\VCKDS
	ļ				1	FDKNYKAPIGADFEMERFEVLGIPF
1092	2442	A	8999	548	811	SSFIKRHILIFEDDWHQTTCCHHPHHP\F*RCQ
1002	2	1.		3.0		FHIFYVSVQNSISPSLSVSSSHPDRPDHEVHQH
		J	J	]		RAAHHHOHGOGPLGHGLVARVG
1093	2443	A	9002	3	2745	ALLGLQQPAQSLILSRSSVMGVRGLQGFVGS
			1	]		TCPHICTVVNFKELAEHHRSKYPGCTPTIVVD
						AMCCLRYWYTPESWICGGQWREYFSALRDF
						VKTFTAAGIKLIFFFDGMVEQDKRDEWVKRR
			1			LKNNREISRIFHYIKSHKEQPGRNMFFIPSGLA
						VFTRFALKTLGQETLCSLQEADYEVASYGLQ
						HNCLGILGEDTDYLIYDTCPYFSISELCLESLD
			Į.			TVMLCREKLCESLGLCVADLPLLACLLGNDII
						PEGMFESFRYKCLSSYTSVKENFDKKGNIILA
						VSDHISKVLYLYQGEKKLEEILPL/VTKQSSFL
						*RNGIISFTRT/INLHGFSKNPKV**LWTNK*YP
						RVQTPNPGKKFPCVQMLNPGKKFPCVQALNP
		1	ŀ	{		GEKFPCIHI/PEPRQEVPTCSDPEPRQEVPTCTG
						PESRREVPMCSDPEPRQEVPMCTGPEPRQEVP MCTGPEARQEVPMCTDSEPRQEVPMCTDSEP
						RQEVPMYTGSEPRQEVPMYTGPESRQEVPMY
1			l			TGPESRQEVLIRTDPESRQEIMCTGHESKQEV
	L		<u> </u>	L	L	TOT POWARA PHY INTERVALENCE TO LEGIS A PARTY OF A

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  PICTDPISKQEDSMCTHAEINQKLPVATDFEFK LEALMCTNPEIKQEDPTNVGPEVKQQVTMVS DTEILKVARTHHVQAESYLVYNIMSSGEIECS NTLEDELDQALPSQAFIYRPIRQRVYSLLLED CQDVTSTCLAVKEWFVYPGNPLRHPDLVRPL QMTIPGGTPSLKILWLNQEPEIQVRRLDTLLA CFNLSSSREELQAVESPFQALCCLLIYLFVQV DTLCLEDLHAFIAQALCLQGKSTSQLVNLQP DYINPRAVQLGSLLVRGLTTLVLVNSACGFP WKTSDFMPWNVFDGKLFHQKYLQSEKGYA VEVL/CRTK*ISAHQIPQPEGSRLQGLHEGEQT
1094	2444	A	9021	97	834	HHWPSPLGLTPRREVGKTGLQLPQDGLWV AREACRAKTDFPGRRFRLWPSCCCRVIVGAE T*HMAEPVSPLKHFVLAKKAITAIFDQLLEFV TEGSHFVEATYKNPELDRIATEDDLVEMQGY KDKLSIIGEVLSRRHMKVAFFGRTSSGKSSVI NAMLWDKVLPSGIGHITNCFLSVEGTDGDKA YLMTEGSDEKKSVKTVNQLAHALHMDKDLK AGCLVRVFWPKAKCALLRDDLVLVDGPGTD VTTELDSWIDKFCTKSSTREITNSGSDT
1095	2445	A	9022	1	537	LVLNSRVEDFVPPEGAGRTLPFALRPLAACW LLHRRARRSSALCPRPRSWGVSGGEGAGARE P*ITSSSCCLSAA/SHLSIQSPNMAGARRIRPQ LAKEKIEGCHICTSVTPGEPQVFLGKDKAFTF DYVFDIDSQQEQIYIQCIEKLIEGCFEGYNATV FAYGQT\GAGKTYTMGTGFD
1096	2446	A	9029	1	285	FFFFNVCKSPKVPKPGCKEESTGTLFKNTLISL GQHSETPSLKKKLAGYSGMCL*SQVLRRLRQ EDCLSPGGGNCRES*SCPYTPAWITERDPV
1097	2447	Λ	9032	716	357	ARSTGFWGEILWCGFLKRSLALSPRVKCSGAI LAHCNFRHAGFPPLSCLSLPNRWEYRRPPARP GKFFLVFLVETGFQC/G*DGLDLLTSRSACLG LPKCWDYRREPAASIIFQTTFFINSK
1098	2448	A	9038	230	652	KVVVMSCEDINISGSFYRNKLKYLAFLCKRTS TNPSQGPYHLWVPSHIFWQTTCGRLPHKTKQ G*AALDHLKVFDRIPLPYDKKKQMAVSATLE VVRPKP*RKFAYLGHWAQKVDWKYQAMTA TMGEKRKVYYQKICYQKK
1099	2449	A	9043	185	372	IIFYSHQQCMRV/WQGCGDIETLIHCW*E*KII HSL/WK/TV*QFLKRLYLHLPHNSVIAFLGISP RKIKTCPQNSCTSMLINAIHNDQKWKKINI
1100	2450	A	9045	763	584	ROSLALSPRLECSGTISAHCRLCPLVFTPLSCL SLTSSWDYRRPPPHPANFLYFK*RRGF
1101	2451	A	9050	275	2	LFFLRKVSNQFLSPSLLPVNFQGFVFAFLLLL FLL/FEMESLPVA/RVECSGTISAHCNLCLPGSS DSPASAS*VAGITDMCRYTQLILFHAS
1102	2452	A	9053	449	1224	KTSMFWKFDLHSSSHIDTLLEREDVTLKELM DEEDVLQECKAQNRKLIEFLLKAECLEDLVSF I\*EEPPQDMDEKIRYKYPNISCELLTSDVSQM NDRLGEDESLLMKLYSFLLNDSPLNPLLASFF SKVLSILISRKPEQIVOFLKKKHDFVDLIIKHIG TSAIMDLLLRLLTCIEPPQPRQDVLN\WFKVQ RNL*HST*NVMDISKYVNLHWGLNKSHSLL* LLLQCVLQWLNEEKIIQRLVEIVHPSQEEDVS SLV
1103	2453	A	9058	403	3	GLHVYDFQVYREHILTLNVKKCSVSFWGLRE WLYLQMYEIIKSPRFPIIKMTDITKCW*GC\GA AGMQI/H/CW\WCVNVGKFWEMS*YYLLKLSI ST/PYDPAIPLLGIYL*ETRVYIHPKTCMRMLIA

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
1104	2454	A	9064	75	393	APFVLAVNC KWLFSSLNITGRGDIIGHLKWLDCR\NCSSFPI KRNRQTHSTESNKLKAGHSFGYN*LIH*NS\V KTDCGCGANSKGVVVVMKV\KTAQQKQTTS YMQIGTTKNSRAT
1105	2455	A	9065	366	778	DLLILRNLAFPELKRRNCISRFYLAYHLHKIYS RSILLCNNCSGFYILSL*QYDVFFFNYFFFRDR AWPCCPGWSAAWLTIVILAHYRRPGLERSCC LSLSSSWDHRRVPPCPANF*/YFSMGFTAFPRL VLNS*TQGI
1106	2456	A	9083	673	816	ESGSLIH* WWENKPAQPLWWEI*QHVQKLPT HFPCDPAIPLLGICPED
1107	2457	A	9086	580	18	KPSSGSFIRAIYIFLSTAHVPALFSVLVRTKLT* AFSQSSVLWAHKQQKTSLSLVIR/ERLQIKTA VRENFLPIRLAKILKLDNVKCWQG/SGSNMSL I/HCWWEYNVIHIIWNSVTFPRKVEHVYITYA PEISVR*IHGGLPTLVHQETHTSVFRGAPSVIP ETR\CRPTKESINKLLHIYTMEHYGDENK
1108	2458	A	9093	540	1	GGNDCSVTPTTEPGRKEIT*KRKF*EKTDRLP GA/PPSRTPPTPYPCPHGDRLLPPSRPLPAGPA SAFPPAERSRGHRRASL*RARWSAAVPRRSA GSASEPVQSRWLRLPVGSDSPPAVPVRVCPAP DSRPAAPGSRLPDPGLDSPAPSRTPSSSVD*GG QRPPPPSGDSLSPPGCCRY
1109	2459	A	9099	1255	1425	HESYHVNPNLCNPVAPTSGAHSIG*KWPSWL GAVAHSCNPSTLVGRGGRITRGQELR
1110	2460	A	9103	242	70	EEQFFFFAVGMFP*VDFLAPASGELWDRLRLT CSRPFTRHQSFGLAFLRVCSSLDSLDDSVVGP SALLSSVL/NQGGRNVLEAREAAKHPTI*RQS LLRKQRNKRMAIP
1111	2461	A	9110	189	121	SFLSVRLECNGAIMAHCALPLPG
1112	2462	A	9113	100	910	RRRGGGSRPRRTPVPAPGPGPSFGMDVRFYP AAAGDPASLDFAQCLGYYGYSKFGNNNNYM NMAEANNAFFAASEQTFHTPSLGDEEFEIPPIT PPPESDPALGMPDVLLPFQALSDPLPSQGSEFT PQFPPQSLDLPSITISRNLVEQDGVLHSSGLHM DQSHTQVSQYRQDPSLIMR\PSST*PDAARSG VMPPAQLTTINQSQLSAQLGLNLGGASMPHT SPSPPASKSATPSPSSSINEEDADEANRAIGEK RAAPDSGKKPKTPKK
1113	2463	A	9120	3452	3051	FLRPSFALVPQAGVQWCALSWLQPPSPRFK*F SCLSLPSSWDYRHVPPRPANFFVLLVETGFLH VGQAGHEPLTSGDPPASASQSAGITGVSHQA WPSFFIFSRDTVLLCCSGWSRTSGLKQSACLS LLKCWDY
1114	2464	A	9122	152	377	NQLPLQQWTFFIYETGFCSVAQAGVQCRDHS SLHP*PPG\SSDPPAPPS*VLGITGQRYHACLII YLYVQTVPQRV
1115	2465	A	9124	553	981	QRPLLRQQLGSWPTCRSLEGDLASPW**RLPG SPRMRRSGT/ATLNLPLSPQGTVRTAVEFQVM TQTQSLSFLLGSSASLDCGFSMAPGLDLISVE WRLQHKGRGRGDLHLPDHHLSVPSSADHPA QQPSQFNGRNLYFLPLFR
1116	2466	A	9135	48 .	410	SASHEPAEHDGGADSLSASQPPRPAGRPAGA QHVHVPPWTDVLAGQDRRAPTAGDGAPWP APGGHVPSTRPHDPAEFHADEAAGRGGRGLQ PAAPHALPAGLPHGPPAPA/PAEGGGTP*GSA GAGGP*GSPAGRACGAAGCRPRPPRPAASSA *NSAGS*GLVEGT*PPGAGHGAPSPAVGARLS

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  CPARTSVQGGTWTC*APAGRPAGLGGWEAE RESAPPSCSAGS*DAD*GAEPWGAGSRSWGS  KSGHWAKECLQPRIPPRPCPICVGPHWKSDCP TCPGAVPRAPGTLPQGSLTDSFPDLLSLVAED *CCLMASEASWTIT\ELWVTLTVEGKSVP/CL NTEATHSTLPSFQGPVSLASITVVGIDGQASKP
	ĺ					LKTPQLWCQLGQYSFMHYFLVIPTCPVPLLG* GILTKLSAFLTIPRLQPHLIAALSPSS
1118	2468	A	9154	471	2	AAGQVVVEVTSHLYLCITSDAAGLRLLPPAES ERGEGGHCPAEAPLPPRPQYCLAKHPLLRKLP EEKIKLDPYLTQHTKINSKQIKYLS/VRAKTTQ LVEGNIGVNLQNTELKQH*INGFLDTTPEAQE TKEKTNKLNFIKKVKRQLAEWEKIFQIA
1119	2470	A	9155	124	207	ACPRLARRRRRVRSLRRRRGWLRARWSRGQ NNMAARRITQETFDAVLQEKAKRYHMDASG EAVSETLQFKAQDLLRAVPRSRAEMYDDVHS DGRYSLSGSVAHSRDAGRESLRSDVFSGPSFR SSNPSISDDSYFRKECGRDLEFSHSNSRDQVIG HRKLGHFRSQDWKFALRGSWEQDFGHPVSQ ESSWSQEYSFGPSAVLGDFGSSRLIEKECLEK ESRDYDVDHPGEADSV/LRGGSQVQARGRAL NIVDQEGSLLGKGETQGLLTAKGGVGKLVTL RNVSTKKIPTVNRITPKTQGTNQIQKNTPSPD VTLGTNPGTEDIQFPIQKIPLGLDLKNLRLPRR KMSFDIIDKSDVFSRFGIEIIK WAGFHTIKDDIK FSQLFQTLFELETETCAKMLASFKCSLKPEHR DFCFFTIKFLKHSALKTPRVDNEFLNMLLDKG AVKTKNCFFEIIKPFDKYIMRLQDRLLKSVTP LLMACNAYELSVKMKTLSNPLDLALALETTN SLCRKSLALLGQTFSLASSFRQEKIL*AVGLQ DIAPSPAAFPNFEDSTLFGREYIDHLKAWLVS SGCPLQVKKAEPEPMREEEKMIPPTKPEIQAK APSSLSDAVPQRADHRVVGTIDQLVKRVIEGS LSPKERTLLKEDPAY WFLSDENSLEYKYYKL KLAEMQRMSENLRGADQKPTSADCAVRAML YSRAVRNLKKKLLP\WQRRGLLRAQG\LRG\ WKARRA\TTGTQTLLFLRAPGLKHHGRQAPG LSQAKPSLPDRNDAAKDCPPDPVGPSPQDPSL EASGPSPKPAGVDISEAPQTSSPCPSADIDMKT METAEKLARFVAQVGPEIEQFSIENSTDNPDL WFLHDQNSSAFKFYRKKVFELCPSICFTSSPH NLHTGGGDTTGSQESPVDLMEGEAEFEDEPP PREAELESPEVMPEEEDEDDEDGGEEAPAPG GAGKSEGSTPADGLPGEAAEDDLAGAPALSQ ASSGTCFPRKRISSKSLKVGMIPAPKRVCLIQE PKGECPPVGTVASSTVLGWWAVRVRDRWR HFNPKEFCAPLQNVSRHSCFPVV
1121	2471	A	9166	272	523	PPRACRPCPRACPCPPT*KCSQPVSWPC PMSSLQGCFYTFKCIIFKGIFLLLISNLIAF**EK
						V/CSHITDSLKFIGKGWVGMVTHACNPGTLG
1122	2472	С	9170	442	236	G*GGWIA*VREFETSLGNM MNRRRFLRPADCIISGMRGTENGACSEGESQI HCGAGGEGVQLVHVVNQPENGCLQFDSTHIT FSKRQN*
1123	2473	A	9171	10	423	MVDRSPLLTSVIIFYLAIGAAIFEVLEEPHWKE AKKNYYTQKLHLLKEFPCLGQEGLDKILEVV SDAAGQGVAITGNQTFNNWNWPNAMIFAAT VITTIGYGNVASKTPGGRLFCGFYGLFGVPFC LTWINALGKFFG

No. of   No. of   No. of   nucleotide (contide conti	SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
Decide   pepide   corde   corresponding   co							D=Aspartic Acid, E=Glutamic Acid,
Sequence	nucl-	peptide		in			
Sequence	eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
mainto acid residue of sequence   residue of sequence   sequence	seq-	uence	1	09/496	correspondi		M=Methionine, N=Asparagine, P=Proline,
Part	uence			914		acid residue	
					amino acid		
1124   2474   A   9173   3   374   Seguence	Ì		ł	1		sequence	
1124	İ		İ				
WEDQGGLIGPESILMI.MLILETRINPYNACI.							
TGSLFYLIGVPSFEPVPSCALQELKPRDRISA   IAHRGGRIEDPPSTILGAR/QGS-WSNRR   IAHRGGRIEDPPSTILGAR/QGS-WSNRR   IAHRGGRIEDPPSTILGAR/QGS-WSNRR   ISSSCILIFQCFQGIHVQKLTLQARPTLFSWWL   CSKPPKETGELERARSGGGDGRGGKQDNV   AWWRRMQKGDFPWDDEDPQSGPFGGGA   LPMGFFYLYFDPGDEITWRIFVQYYLARGL   VDRLEVVNKQSVRVIPAPGTSSEVRGEFKAE   YCHKFISCKNVYFYFQ   VDRLEVVNKQSVRVIPAPGTSSEVRGEFKAE   YCHKFISCKNVYFYFQ   VDRLEVVNKQSVRVIPAPGTSSEVRGEFKAE   YCHKFISCKNVYFYFQ   VDRLEVVNKQSVRVIPAPGTSSEVRGEFKAE   YCHKFISCKNVYFYKQ   VDRLEVVNKQSVRVIPAPGTSSEVRGEFKAE   YCHKFISCKNVYFSKSRAGLGARKLKAYRTM   EYMAESTDRSFGHILCCECGYPISVPAQYCV   ACKRSFFIVHYCFKLFHPFSTSSAFITSHY   LTFTSTIS   VLKFILLGTMDEQSQGMQGPPVPQFQPQKAL   RCHARLSFFILVRKLIGRGQFSEVYRAAC   LLDGVPVALKKVQFFDLMADARARADCIKEID   LLKQLMIPNVKYYASFIEDNELNIVLELADA   GDISKNIKHFKKQGKLIPERTVYFYQLCS   ALEHMISRRVMIRDIRPANVFITATGVYKLG   DLGGFFSSKTTAAHISLVGTYYYMSPERIEID   NG   GTSWKIPSAAVSESSPNGAAYASGLPCGVRG   PPWAGLALIPSPTLMALLRRFTVSSDLDNIDT   RATTKJRVVATITTRARIEDMRATALTRPD   ATTAQIPKLPVTTVCNRRANFGIPPSVL   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKKQCKLIPETGV   LALLGYLAVRFPLPKKQCKDSLINLKICKEN   PLAFLALELLFAIRAFATGGGYSGGLRLSVD   CVNKTESNLSIDAFAYPTRILITYCCAYDV   LALLGYLAVRFPLPKKQCK   LALLGYLAVRFPLPKKQCCH   LALLGYLAVRFPLPKKQCCH   LALLGYLAVRFPLPKKQCCH   LALLGYLAVRFPLPKKQCCH   LALLGYLAVRFPLPKKQCCH   LALLGYLAVRFPLPKKQCCH   LALLGYLAVRFPLPKVQCA   ESGCISEKEKGVIVLNCSRLLTYTSCAPTU   PAAGLRAVREESPSNLAALTLCYKAYEKUQCA   ESGCISEKEKGVIVLNCSRLLTYTSCAPTU   LEVELPPTCVYERGPTCC   GRERIHALALGSMYLVLLIV   LAVRLEDPTCVYRGARGGGGEDDEHARPLAE   SLLLLAIDLEFPTCVYSTER GWANTANA   RADGAPAPAGEGGGTTLGGNYLAAC   GVPSIVALCYALGTTISKSCAPTU   LAVRLEDPTGVYRGARGGGGTUTLGGNYLALTYL   LKPLPPTCVYRGARGGGTLAVVLALLSWPLHSNS   NVYFVV	1124	2474	A	9173	3	374	GPSPSLLVLLPQEPGGTGTPVRAGAGAGMWL
ILAHRGGRIÐPPENTLGARGGS**WSNRR   188		1					
1125							
CSKPPKETGEL BNABSGGDGGRGGKODNV							
AWWRM.QKG.ODFPW.DEDFPQSGFGGQA	1125	2475	A	9179	704	188	ESSSGLLFQCFQGIHVQKLTLQARPTLFSWWL
LPMGFYLYFRIPGGERTWKHFYOYYLARGL							
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ACLRSSPHIYHCIPKLEHIPFSKTSSSAFITPSHY	1127	24//	A	9185	] 1	321	
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1128							
RPDMGYNTLANFRIEKKIĞRGQU'SEVYRÄAC   LUBGYPVALKKVQIFDLMDAKARADCIKEID   LLKQLNHPNVIK'YYASPIBDMENIV'LELADA   GDLSRMIKHFKKQKRLIPERTVWKYYVQLCS   ALEHMHSRRVMHRIKPANVFITATGVVKLG   DLGLGRFFSSKITAHSLVGTPYYMSPERIID   NG	1100	0.470		0106	100	0.15	
LUDGYPVALKKVQIELMAKARADCIKEID	1128	2478	A	9186	183	847	VLKFLLLQTMDEQSQGMQGPPVPQFQPQKAL
LLKQLNHPNVKYYASPIEDNELNIVLELADA GDLSRMIKHRKQRRIPERTVVKXYFQLCS ALEHMHSRRVMFRDIKPANVFITATGVVKLG DLGLGRFSSKTTAAHSLVGTPYYMSPERHID NG  1129 2479 A 9190 1 370 GTSWKIPSAAVSSESSPNGAAYASGLPCGVRG PPWAGLALLPSPTIMALLRRPTVSSDLDNIDT RATTKIRVVATITRARIEDMRHSATALTRPD ATTAQIPKLPVTTVCNRRANPGIPPSVL  1130 2480 A 9194 131 487 AYLKRLPVPESTIPGARLTVSEWLRLLPFLGV LALLGYLAVRPFLPKKRQQKDSLINLKIQKEN PRVVNENLEDLCTKAAVGCRWRSKTPPAC DGSHNKHNELTGDNVGPLIKKKE  1131 2481 A 9201 184 605 KELVDEKSERGRAMDPVSQLASSGTFRVLKE PLAPIRALELLPAIFAFATCGGYSGGLRLSVD CVNKTESNLSIDIAFAYPFRLHQVTFEGIPTCE GKERHKLALIGDSSSAEFFGTVAGFAFLYSL AATGVYIFFQNKY  1132 2482 A 9206 1 852 GGGRAGAGSRDMGSTDSKLNFRKAVIQLTTK TQPVEATDDAFWQFWADTATSVQDVFALV PAAEIRAVREESPSNLATLCYKAVEKLVQGA ESGCHSEKEKQIVLNCSRLLTRVLPYIEEDPD WRGFFWSTVPGAGRGQGEEDDEHARPLAB SLLLAIADLLFCPDFTVQSHRRSTVDSAEDVH SLDSCEYIWAGAGYGFAHSPQPNYHDMNRME LLKLLITCTSEAMYLPPAFESWQHKTHWFSS PVSSENRHALPLFTSLLNTVCAYDPVEYGIPY NHLY  1133 2483 A 9208 1165 1463 GPRARVQGFSGADIVKFMALGSMYLVLTLIV AKVLGAEPCCGPLKNRVLRPCPLP/VPLPPP HPQPSRGNPVGCLPTYKVVYKLLSWPLHSNS NVYFIV  1134 2484 A 9210 66 1586 MAGAGPKRRALSAPVAEKEEAREKIMAAK RADGAAPAGEGGGYTLLGKGVAVIVV ALMGSGIFVTIFTGVLKEAGSPGLALVWWAAC GVFSIVGALCYAELGTTISKSGGDYAYMLDV YGSLPAFLKLUELIRPSSQVIVALVATYL LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAKLLALALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVSNIVALVFSG							
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ALBEMHSRRVMHRDIKPANVFITATGVVKLG   DLGLGRFFSSKTTAAHSLVGTPYYMSPERHID   NG							
DLGLGRFFSSKTTAAHSLVGTPYYMSPERIHD   NG							
1129   2479   A   9190   1   370   GTSWKIPSAAVSESSPNGAAYASGLPCGVRG PPWAGLALLPSPTLMALLRRPTVSSDLDNIDT RATTKIRVVATITRARIEDMRHSATALTRPD ATTAQIPKLPVTTVCNRRANPGIPPSVL ATTAQIPKLPVTTVCNRRANPGIPPSVL ATTAQIPKLPVTTVCNRRANPGIPPSVL ATTAQIPKLPVTTVCNRRANPGIPPSVL ALLGYLAVRPFLPKKKQQKDSLINLKIQKEN PKVVNEIDELCITKAAYCRCWRSKTPPAC DGSHNKHNELTGDNVGPLILKKKE PACADGSHNKHNELTGDNVGPLILKKKE PACADGSHNKHNELTGDNVGPLILKKKE PACADGSHNKHNELTGDNVGPLILKKKE PLAFIRALELLPAIFAFATCGGYSGGLRLSVD CVNKTESNLSIDIAFAYPRILHQVTFEGIPTCE GKERHKLALIGDSSSAEFFGTVAGFAFLYSL AATGVYIFFQNKY ATTAGVATATSVQDFALV PAAEIRAVREESPSNLATLCYKAVEKLVQGA ESGCHSEKEKQIVLNCSRLLTRVLPYIFEDPD WRGFFWSTYPGAGRGGGGEDDEHARPLAE SLLLAIADLLFCPDFTVQSHRRSTVDSAEDVH SLDSCEYIWEAGVGFAHSPQNYHIDMNRME LLKLLLTCFSEAMYLPPAPESWQHRTHWFSS FVSSENRHALPLFTSLLNTVCAYDPVEYGIPY NHLY SLDSCEYIWEAGVGFAHSPQNYLPTHMPSS FVSSENRHALPLFTSLLNTVCAYDPVEYGIPY NHLY SLDSCEYIWEAGVGFAGGGCDEDDEHARPLAE SLLKLLLTCFSEAMYLPPAPESWQHRTHWFSS FVSSENRHALPLFTSLLNTVCAYDPVEYGIPY NHLY SLDSCEYIWEAGVGFAHSPQNYLPLAFUNG AKVLRGAEPCCGPLKNRVLRPCPLPVPLPPP HPQPSRGNPVGCLPTYKVVYKLLSWPLHSNS NVYFIV AKVLRGAEPCCGPLKNRVLRPCPLPVPLPPP HPQPSRGNPVGCLPTYKVVYKLLSWPLHSNS NVYFIV LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAKLLALALILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG				1			
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RATTKIRVVATITRARIEDMRHSATALTRPD   ATTAQIPKLPVTTVCNRRANPGIPPSVL	1129	24/9	A	9190	1	370	
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SLLLAIADLLFCPDFTVQSHRRSTVDSAEDVH SLDSCEYIWEAGVGFAHSPQPNYIHDMNRME LLKLLTCFSEAMYLPPAPESWQH/RTHWFSS FVSSENRHALPLFTSLLNTVCAYDPVEYGIPY NHLY  1133 2483 A 9208 1165 1463 GPRARVQGFSGADIVKFMALGSMYLVLTLIV AKVLRGAEPCCGPLKNRVLRPCPLP/VPLPPP HPQPSRGNPVGCLPTYKVVYKLLSWPLHSNS NVYFIV  1134 2484 A 9210 66 1586 MAGAGPKRRALSAPVAEEKEEAREKIMAAK RADGAAPAGEGEGVTLQGNITLLKGVAVIVV AIMGSGIFVTPTGVLKEAGSPGLALVVWAAC GVFSIVGALCYAELGTTISKSGGDYAYMLDV YGSLPAFLKLWIELLIIRPSSQYIVALVFATYL LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAAKLLALALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG	1			1			
SLDSCEYIWEAGVGFAHSPQPNYIHDMNRME LLKLLTCFSEAMYLPPAPESWQH/RTHWFSS FVSSENRHALPLFTSLLNTVCAYDPVEYGIPY NHLY  1133 2483 A 9208 1165 1463 GPRARVQGFSGADIVKFMALGSMYLVLTLIV AKVLRGAEPCCGPLKNRVLRPCPLP/VPLPPP HPQPSRGNPVGCLPTYKVVYKLLSWPLHSNS NVYFIV  1134 2484 A 9210 66 1586 MAGAGPKRRALSAPVAEEKEEAREKIMAAK RADGAAPAGEGEGVTLQGNITLLKGVAVIVV AIMGSGIFVTPTGVLKEAGSPGLALVVWAAC GVFSIVGALCYAELGTTISKSGGDYAYMLDV YGSLPAFLKLWIELLIIRPSSQYIVALVFATYL LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAAKLLALALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG							,
LLKLLTCFSEAMYLPPAPESWQH/RTHWFSS FVSSENRHALPLFTSLLNTVCAYDPVEYGIPY NHLY  1133 2483 A 9208 1165 1463 GPRARVQGFSGADIVKFMALGSMYLVLTLIV AKVLRGAEPCCGPLKNRVLRPCPLP/VPLPPP HPQPSRGNPVGCLPTYKVVYKLLSWPLHSNS NVYFIV  1134 2484 A 9210 66 1586 MAGAGPKRRALSAPVAEEKEEAREKIMAAK RADGAAPAGEGEGVTLQGNITLLKGVAVIVV AIMGSGIFVTPTGVLKEAGSPGLALVVWAAC GVFSIVGALCYAELGTTISKSGGDYAYMLDV YGSLPAFLKLWIELLIIRPSSQYIVALVFATYL LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAAKLLALALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG							
FVSSENRHALPLFTSLLNTVCAYDPVEYGIPY   NHLY							
NHLY	1					ľ	
1133   2483   A   9208   1165   1463   GPRARVQGFSGADIVKFMALGSMYLVLTLIV AKVLRGAEPCCGPLKNRVLRPCPLP/VPLPPP HPQPSRGNPVGCLPTYKVVYKLLSWPLHSNS NVYFIV     1134   2484   A   9210   66   1586   MAGAGPKRALSAPVAEEKEEAREKIMAAK RADGAAPAGEGEGVTLQGNITLLKGVAVIVV AIMGSGIFVTPTGVLKEAGSPGLALVVWAAC GVFSIVGALCYAELGTTISKSGGDYAYMLDV YGSLPAFLKLWIELLIIRPSSQYIVALVFATYL LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAAKLLALALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG							
AKVLRGAEPCCGPLKNRVLRPCPLP/VPLPPP HPQPSRGNPVGCLPTYKVVYKLLSWPLHSNS NVYFIV  1134 2484 A 9210 66 1586 MAGAGPKRRALSAPVAEEKEEAREKIMAAK RADGAAPAGEGEGVTLQGNITLLKGVAVIVV AIMGSGIFVTPTGVLKEAGSPGLALVVWAAC GVFSIVGALCYAELGTTISKSGGDYAYMLDV YGSLPAFLKLWIELLIIRPSSQYIVALVFATYL LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAAKLLALALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG	1133	2483	A	9208	1165	1463	
HPQPSRGNPVGCLPTYKVVYKLLSWPLHSNS NVYFIV  1134 2484 A 9210 66 1586 MAGAGPKRRALSAPVAEEKEEAREKIMAAK RADGAAPAGEGEGVTLQGNITLLKGVAVIVV AIMGSGIFVTPTGVLKEAGSPGLALVVWAAC GVFSIVGALCYAELGTTISKSGGDYAYMLDV YGSLPAFLKLWIELLIIRPSSQYIVALVFATYL LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAAKLLALALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG		]					
NVÝFIV  1134 2484 A 9210 66 1586 MAGAGPKRRALSAPVAEEKEEAREKIMAAK RADGAAPAGEGEGVTLQGNITLLKGVAVIVV AIMGSGIFVTPTGVLKEAGSPGLALVVWAAC GVFSIVGALCYAELGTTISKSGGDYAYMLDV YGSLPAFLKLWIELLIIRPSSQYIVALVFATYL LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAAKLLALAIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG							
RADGAAPAGEGEGVTLQGNITLLKGVAVIVV AIMGSGIFVTPTGVLKEAGSPGLALVVWAAC GVFSIVGALCYAELGTTISKSGGDYAYMLDV YGSLPAFLKLWIELLIIRPSSQYIVALVFATYL LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAAKLLALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG	L						,
RADGAAPAGEGEGVTLQGNITLLKGVAVIVV AIMGSGIFVTPTGVLKEAGSPGLALVVWAAC GVFSIVGALCYAELGTTISKSGGDYAYMLDV YGSLPAFLKLWIELLIIRPSSQYIVALVFATYL LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAAKLLALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG	1134	2484	A	9210	66	1586	MAGAGPKRRALSAPVAEEKEEAREKIMAAK
AIMGSGIFVTPTGVLKEAGSPGLALVVWAAC GVFSIVGALCYAELGTTISKSGGDYAYMLDV YGSLPAFLKLWIELLIIRPSSQYIVALVFATYL LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAAKLLALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG	1						
GVFSIVGALCYAELGTTISKSGGDYAYMLDV YGSLPAFLKLWIELLIIRPSSQYIVALVFATYL LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAAKLLALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG							
YGSLPAFLKLWIELLIIRPSSQYIVALVFATYL LKPLFPTCPVPEEAAKLVACLCVLLLTAVNC YSVKAATRVQDAFAAAKLLALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG							
LKPLFPTCPVPEEAAKLVACĽCVLLLTAVNC YSVKAATRVQDAFAAAKLLALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG							
YSVKAATRVQDAFAAAKLLALIILLGFVQI GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG	1						
GKGDVSNLDPNFSFEGTKLDVGNIVLALYSG							YSVKAATRVQDAFAAAKLLALALIILLGFVQI
LFAYGGWNYLNFVTEEMINPYRNLPLAIIISLP							
							LFAYGGWNYLNFVTEEMINPYRNLPLAIIISLP

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion IVTLVYVLTNLAYFTTLSTEQMLSSEAVAVDF GNYHLGVMSWIIPVFVGLSCFGSVNGSLFTSS RLFFVGSREGHLPSILSMIHPQLLTPVPSLVFT CVMTLFYAFSKDIFSVINFFSFFNWLCVALAII GMIWLRHRKPELERPIKVNLALPVFFILACLF LIAVSFWKTTPWSVASDFTIILSGLPVYFFGV WWKNKPKWAPPGHLSPRPSCVRSSCMVVPQ
1135	2485	A	9216	40	410	RDRLPPAYFCRPVVCVVTALDVG\SPESQEM DLVAFEDVAVNFTQEEWSLLDPSQKNLYREV MQETLRNLASIGEKWKDQNIEDQYKNPRNNL RSLLGERVDENTEENHCGETSSQIPDDTLNK
1136	2486	A	9223	3	983	RRRRSRYRRCSRFPRPGPLAVSMPHAFKPG DLVFAKMKGYPHWPARIDDIADGAVKPPPN KYPIFFFGTHETAFLGPKDLFPYDKCKDKYGK PNKRKGFNEGLWEIQNNPHASYSAPPPVSSSD SEAPEANPADGSDADEDDEG\RGVMAVTAVT ATAASDRMESDSDSDKSSDNSGLKRKTPALK MSVSKRARKASSDLDQASVSPSEEENSESSSE SEKTSDQDFTPEKKAAVRAPRRGPLGGRKKK APSASDSDSKADSDGAKPEPVAMARSASSSSS SSSSSDSDVSVKKPPRGRKPAEKPLPKPRGRK PKPERPPSSSSSD
1137	2487	A	9229	21	239	LFPRLECRDPVTVNCTLNLPGSKNAPTTASQV GSTWNYRGGLPHPTNFFVKTGFRCSQAGLKL RGSREPPAWA
1138	2488	A	9231	207	443	TRSVGVNTCEVGVVTEPECLGPCEPGTSVNL EGIVWHETEEGVLVVNVTWRNKTYVGTLLD CTKHDWAPPRFCESPTSDLEMRGGRGRGKR ARSAAAAPGSEASFTESRGLQNKNRGGANGK GRRGSLNASGRRTPPNCAAEDIKASPSSTTNKR KNKPPMELDLNSSSEDNKPGKRVRTNSRSTP TTPQGKPETTFLDQGCSSPVLIDCPHPNCNKK YKHINGLRYHQAHAHLDPENKLEFEPDSEDK ISDCEEGLSNVALECSEPSTSVSAYDQLKAPA SPGAGNPPGTPKGKRELMSNGPGSIIGAKAGK NSGKKKGLNNELNNLPVISNMTAALDSCSAA DGSLAAEMPKLEAEGLIDKKNLGDKEKGKK ANNCKTDKNPSKLKSARPIAPAPAPTPPQLIA IPTATFTTTTGTIPGLPSLTTTVVQATPKSPPL KPIQPKPTIMGEPITVNPALVSLKDKKKKEKR KLKDKEGKETGSPKMDAKLGKLEDSKGASK DLPGHFLKDHLNKNEGLANGLSESQESRMAS IKAEADKVYTFTDNAPSPSIGS TRRGQPWRRAAAAGILPGREAAACLPSC/AS
						VTAAVSGLLVGYELGIISGALLQIKTLLALSC HEQEMGVSSLVIGALL
1140	2490	A	9238	248	328	MAQGNNYGQTSNGVADESPNMLVYRKV
1141	2491	A	9242	2	535	FVEAAVKMLGSLVLRRKALAPRLLLRLRSP TLRGHGGASGRNVTTGSLGEPQWLRVATGG RPGTSPALFSGRGAATGGRQGGRFDTKCLAA ATWGRLPGPEETLPGQDSWNGVPSRAGLGM\ WPWAAALVVHCYSKSPSNKDAALLEAARAQ \NMQEVSRNRCALLHSAAVQEYGYGN
1142	2492	A	9245	157	466	HILCFWFFVGLFLPEQQIMLFATLLRMAQGCD FALGNDFLNITTKAQA/TKEKLDKLDFIKIKTC CTSMDAIEKTEPLTKWTKAFVSHVSYKRLLF GICKEYSRQ
1143	2493	A	9247	264	115	GLPQQTSTIQPPGTPDGARDFTSTIQPPGAPDG ARDSTSIIRMGPEIPPP

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
1144	2494	A	9260	1	401	KKVPGRLSEMSFSLNFTLPANTTSSPVT\DCGP SLGLAAGIPLLVATALLVALLFTLIHRRRSSIE AMEESDRPCEISEIDDNPKISENPRRSPTHEKN TMGAQEAHIYVKTVAGSEEPVHDRYRPTIEM ERRR
1145	2495	A .	9264	175	411	METIWIYQFRLIEIGDSTVGKSCLLHRFTQGRF PGLRSPACDPTVGVDFFSRLLEIEPGKRIKLLL WDTAGQERFISIT
1146	2496	A	9277	592	814	MFTYLEGREGIKSQPKMEPHSVT\RLECSGMI SAHCSLNLPGTSDSPASASR/VAGTTGMRHHA WLIFAFLVETGF
1147	2497	A	9279	1255	2	FRRGRRGEEEKEEEEEEGWVNGMENSHPP HHHHQQPPPQPGPSGERRNHHWRSYKLMIDP ALKKGHHKLYRYDGQHFSLAMSSNRPVEIVE DPRVVGIWTKNKE\LELSVPKFKIDEFYVDQV PPKQVTFAKLNDNIRENFLRDMCKKYGEVEE VEILYNPKTKKHLGIAKVVFATVRGAKDAVQ HLHSTSVMGNIIHVELDTKGETRMRFYELLV TGRYTPQTLPVGELDAVSPIVNETLQLSDALK RLKDGGLSAGCGSGSSSVTPNSGGTPFSQDTA YSSCRLDTPNSYG/QGTPLTPRLGTPFSQDSSY SSRQPTPSYLFSQDPAVTFKARRHESKFTDAY NRRHEHHYVHNSPAVTAVAGATAAFRGSSD LPFGTVGGTGGSSGPPFKAQPQDSATFAHTPP PAQATPAPGFR
1148	2498	A	9302	1026	6	IASIQNADTMPGVGLLVSHFSTLVSRQRCPNY ADPQNLTDVSIFLLLEVSGDPELQPVLAGLFL SMCLVT.VLGNLLIILAISPDSHLHTPMYFFFSN LSLPDV\GFTSTTVPK\MIVDI\QSRSRVISYAG CLTQKSLFAIFGGTEE\NMLLSVMAYDRFVAI CHPLYHSAIMNPCFCAFLVLLSFFFLSLLDSQL HSWIVLQFTIIKNVEISNFVCDPSQLLKFACSD SIINSIFIYFHKDPERQLVLAGLFLSMCLVTVL GNLIIILDVSPDSHLPTPMYFFLSNLSLPDIGFT STTVPKMIVDIQSHGRVIFYAGCLTQMSLFAIF GGMEERHAPECDGL
1149	2499	A	9303	1	699	MASQEKDIFIGWGTIHLFRKPQRSFFGKLLRE FRLVAADRSMGRYMLFGVINLICTGFLLMWC SSTNSIALT\SYTYLTIFDLFSLMTCLISYWVTL RKPSPVYSFGFERLEVLAVFASTVLAQLGALF ILKESAERFLEQPEIHTGRLLVGTFVALCFNLF TMLSIRNKPFAYVSEAASTSWLQEHVADLSR SLCGIIPGLSSIFLPRMNPFVLIDLAGAFALCIT YMLIEI
1150	2500	A	9308	797	693	DRSTSVTRAGVQWCSLGSLQPRTPGLLRSSCL SLP
1151	2501	A	9309	205	406	VAIKELPVLWKWSKPTR\TAKEPPQTQQRAG SKTAAPPCQWSRMASEGPNIPCPGARHSDKQ FLICTI
1152	2502	A	9314	913	504	KPSPLITPPAVVLPPSAVLNLVNTFSSFPQVEV QGPLCGPRKGRLAVTIPFFGLS/LPKYMDHRR PPPHR\EIFFVFLAETGFHRASQAGPDLPTS/S/I PPTSA/FPKCWEYRSEPQCLPGCLSFSGILLDL GTNVSLRAA
1153	2503	A	9315	392	1	HPHRPRPGFRSPARSSRPCPVLTSLLPPFPSPSP PADDLVKAGRDRKDPQVR/ERRLRPNPGRLG GPR\PRPARARS/CHQPRLTRVCPRSPPPEARA PAPAAPARGRGAPKRNRPRTDTRAPRGSSAR PGNS

1154	SENNVL GGGPAK
1156	GGGPAK
EAAQRHCRASVSILRMRRPGQGSSRF RGPDSHRLREPPPSPP  1157 2507 A 9327 152 292 YERRGRSQGGSHPAGAQPGGRAIG. KEPLWEGLQRSGSPLPG  1158 2508 A 9328 1 430 QELKQGPNPLAPSPSAPSTSAGLGDC LSKTFSVSSALAMLQERRCLYVVLTI	
KEPLWEGLQRSGSPLPG   1158   2508   A   9328   1   430   QELKQGPNPLAPSPSAPSTSAGLGDC   LSKTFSVSSALAMLQERRCLYVVLTI	
LSKTFSVSSALAMLQERRCLYVVLTI	AGWQS
SESGLLVSCFDIGNLVVVVFVSYFRG RVAAVGGLLDLEGGEMI	OSRCFL RYSLKS
1159 2509 A 9334 108 383 KGNQVNGNGNQLKRKHESMCPVSL' LMEAGLPQKQAERADELFEAGLVIY' VLNAL\YSSVGLQWFKESDLSHLRLL	VKLDER
1160 2510 A 9338 2 430 FVGRPRGLSDRLEDLFLAGFRVGERL KRYVRILLLGEGAEHVADPVPGGRG DHTDQELREEIHKANVERVVHDVSQ RTKWIPLV/RWGDHA/EGPVGIKSYLI EAELPIMSQLTEIETCVEC	RTAAM VPRGEA EATIEKI
1161 2511 A 9341 1 390 NSRVDDFVAPGLSEAGKLLGLEFPER AVG/CSPMSGVISMSAPFFLGKIIDAIY DYSDNLTRLCLGLSGVFLCGAAANA QTSRQRVVKRLRTSLFSSILGQEVAFS GELI	TNPTV IRVYLM
1162 2512 A 9343 84 837 QGRFRAFCWQRDFLQPPGMRLSALL TLPPHYRYGMSPPGSVADKRKNPPW VVEPISDEDWYLFCGDTVEILEGKDA VVQVIRQRNWVVVGGLNTHYRYIGK GTMIPSEAPLLHRQVKLVDPMDRKPI FTEAGERVRVSTRSGRIIPKPEFPRAD WIDGPKDTSVEDALERTYVPCLKTLC AMGIKETR\NTRRSIGIEPGAEQLLPNI G	IRRRPV IGKQGK CTMDYR FEIEWR GIVPET QEEVME
1163 2513 A 9346 967 616 DSLALSPRLECSGAISAHCNLTPPGFT LPSSWAYRCASPHPDNFFVFLVESGF AGLKLLISSDPPTSA/FPKCWDYRRDW FSSYQRNNPDLILNDTIMPNIK	HHVGQ
1164 2514 A 9347 3 1099 SSFPTCMRTVFHSNTSVSSLLHRPGHY HGGWRHHRDHTAIDEWDFNPSKFLF FSVLLPLRLDGIIQWSYWAVFAPIWL VAGASVGAGVWARNPRYRTEGEAC MLIAVGIHLLLLMFEVLVCDRVERGT VFMPLFFVSPVSVAACVWGFRHDRS SVNILQFIFIALKLDRIIHWPWLVVFVI SFLCLVVLYYIVWSLLFLRSLDVVAE VTMAISWITIVVPLLTFEVLLVHRLDG YVSIFVPLWLSLLTLMATTFRRKGGN AIRRDF/CQDQLPQPTGKPPPPPLTDHI LPLQNKDRGSWPASRGSPRLL	YTCLLL WKLLV VEFKA THFWLL LELEILC PLWILM QRRTH SHNTFS THWWF
1165 2515 A 9362 547 991 DVSIGPPLLRRPCSGREQTRSLSFPSDI VPEGVRLADGPGHCKGRVEVKHQNG CQTGWSLRAAKVVCRQLRCGRAVLT TKHAYGRKPIWLSQMACSGPEPTLHI LGEDTLFHVEYTSVHGRERLSAKD	QWYTV I\QKRC
1166 2516 A 9363 201 387 PPILRWTPPSGKNFFFFFFESEFY/SSF	
	VAN

SEQ ID NO: of nucl- cotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  PRPLILYAPAP\PAGTAFIPHSHPPPPDLLRPT
						ATPA/TPCPSLPPPPRPLHPTQPSTALLPDPPPW PLPFPPPSS/RPPRPDCSTSYSPTFPPPT
1168	2518	A	9375	511	15	MMLSEETSAVRPQKQTRFNGAKLVWMLKGS PITVTSAVIIVLMLLMM/IFSPWLATHDPNAID LTARLLPPSAAHWFGTDEVGRDLFSRVLVGS QQSILAGLVVVATTGMIGSPLECLFGELGGRA DAIFMRVMDIMRS/IPSLVLTMEKTAALGPSL FNAMQASSEH
1169	2519	A	9377	42	410	GNGRVAPRDPGAVASAEPGLTTHDSGVNPN NSARRMEAMASGSNWLSGVNVVLVMAYWS LVFVLLFIFAKRQIMRFAMKSLRGPHGPVGH NAPKDLKEEIDILLSRVHNIKYEP\HLLADDDA
1170	2520	A	9378	302	1303	GVSGFSASVLRQRRMEDELEPSLRPRTQIQGR ILLTICAAGIGGTFQFGYNLSIINAPTLHIQEF TNETWQARTGEPLPDHLVLLMWSLIVSLYPL GGLFGALLAGPLAITLGRKKSLL\VNNIFVVS AAILFGFSRKAGSFEMIMLGRLASWGVNAGV SMNIQP\MLPGGESAPKELRGAVAMSSAIFTA LGIVMGQVVGLSTTAATGLRGL\AGELEELEE ERAACQGCRARRPWELFQHRALRRQVTSLV VLGSAMELCGNDSVYAYASSVFRKAGVPEA KIQYAIIGTGSCELLTAVVSVSLEGALPPPAL WGGTPRSFALNQFTLQKKKK
1171	2521	A	9381	2	412	RGPASAQEDERARTAPLERVRARGRMTTSSA LFPSLLPCSWSTSNKYLAEFRAGKMSLKGTTE TPDKRKGLAY/IQQTDDSLIHFCWKDRTSGNV EDDLIIFPDDCEFKRLPQCPNGRVYVLKFKAG SKRLFFWMQEP
1172	2522	A	9384	20	355	GWNGRSTEASPAAEAPHVPHKET\KAAMGTQ CTHGGKVRPDPHDMLTTVVHKIKLFVLCHSL LQLCAIMISDYLKSSIYTVEKRLGLFRPTSGLL ASFNEVGNTALIVLESY
1173	2523	A	9393	430	87	LCQCIVPGQQKETFSLNPSSATVRFYL*LSLQ QRKEDQ*IIL*YHLNKDCLHIFMSAITLYMKI* KIFVLFDFNIMFETPFYII*FIFLFSQNLKRIRQV IRPPISFSKINNGP
1174	2524	A	9397	77	374	ERLEIGRLGGERGSGPASCLRVIDVSGMWDQ RLVKLALLQLLRAFYGIKVKGVRVHRDCGTF ESSSTLIRVS*FGVPCNALAHFGVTHF*YILDF LGML
1175	2525	A	9399	66	397	HESSRADRDKMDTRGSTYTDADPVNKSGGT AKMNKWSKGKVRDKLNNLVLFDTATYDKL CKEVPNYKLITLAVVSERLKIPGSLARAALHE LLSRGLI*LVIQHIAQVIY
1176	2526	A	9408	2	299	LDLTHVLSLSISLTVTILGTTFGMVIPLLDVVY GERGYAQNGDF*DAQLDDYSFSCYSHAQVN GAPNSLTRAYDDP*VKISGLECQKVGALVEV KCLNL
1177	2527	A	9416	2	402	CNFLRSSRIRVHSTPAASTMPPKVDPNEIKVV YLRCTGGEVRATSALAPKIGPLGLSSIKVGVD FV*ATGDWNVL1ISVILTIRILLSHIFVVPPFFCF DHLIAFWDLQSLIFLHVIFSLFITLLLFCFFSIF
1178	2528	A	9419	142	426	TPLFDLWPRVVLSWLETVLTSLRTRRAASGPP ACRIMPTTVDDVLEHGGEVHFLQKQMLYLL ALI*DTFAPIYVGIVFLGFTPDHRCRSPGVAEL
1179	2529	A	9420	1450	1655	LSSAGTKMNLN*KNYWPGASAHACNPSTLG GQSRCITRSGDRDHPG*HGETPSVLKIQKISRA WWRAP

SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nuclcotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide		in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence			914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
	ļ			amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
	j	J	J	peptide	_	/=possible nucleotide deletion, \=possible
				sequence		nuclcotide insertion
1180	2530	A	9422	176	375	HRPQTTRPDWKPRT*PQGK*GRLSSEISPASPP
			1			SRFSRSTKPVPPKADPPARQKLTGVLHAPLLK
						L
1181	2531	A	9436	2	274	PIAASLRMYNLOPYTEENLICTAFATMVETVP
	J					IARTILDRLTGIPHGYCFVE*ADWATADKCVH
						IYNGKPLPGATPLLSLQLHQLAHLGS
1182	2532	A	9442	3	240	VDKCSSKSIVLSEYCPHCMCSLSTDPKPFGQL
						SMILK*MGAGDEKISAMGKARVDHRELYLGL
						LYPTEDYKLTFRARH
1183	2533	A	9444	384	3	LKDFQPWALHDWPLFCCCTFLLFLVLECFTR
						KGCSGWAPWLSLQCQHFGRPRWADHLRSGV
						RDQPGQYSKTTFLPKIQKLAGHSGAHL*S*LL
	1					ERMRWKNRLNPGGRSCSEPRWHHCTPGWAT
1						ERG
1184	2534	A	9462	391	655	LSGFKSLMPKIPLQYIYVRVRTTWSFCLPLDG
			1			RKLMLS*YSK*LT*KYNILPEYSRMTLPPGMV
						IHTCNPSTLGGRAGWIV*AQEFET
1185	2535	A	9467	215	566	RCPMWQGQASRMDPAKAKDREASTCCSLA
						WWWGWECWVRALKLSSGPAGPLACWVAK
		į				KKSLSLSGPVYPSEKGAGLYVF*DRVSLCHPG
						WSAVVQFWLTAASNSCFSLLSSWDYRCA
1186	2536	A	9468	275	452	HIPQLHTKTHYVPTRMVNKI*QIDNSKPWQR
1100		**				GG*TGILTHCW*ESKLVQPLWKIVWHYQ
1187	2537	A	9469	388	3	EVAPGPSQILPRRVTDGGDRPQFSLPGPRLPQ
110.			7.03	500	-	SSRGAEPCLSNCIHSPAPRKQRMGDSDQ*STP
						NPASPHPEAPQEPWDSASGSVGSFSLGRGAK
						ASS*VPGKGRGPRQGSELLAETILELFLALAN
	ļ		1			S
1188	2538	A	9471	124	397	TMDKKNRHGNSLDMASEIHMTGPMCLIENTT
						GRLMANPEALKILSAITQPMVEEAIAGLYRAC
			1			*FYLTNNLAGMKKGLCLGSTEQAHTIGI
1189	2539	A	9480	584	769	GHVQSQHFGRPRRADHLRSGDRDHPG*HDET
						PSLLKIQKISWAWWRAPVVPATWEAEAEEW
1	}	ļ	l			R
1190	2540	A	9483	463	86	VTVGLTLLLRGAPRFTAG*PPSGGGPPLAPLL
						PRQHCTLQTHRHLHPEAPVKV*KT*RLFPGLR
				İ		GASSCRRRRCNPVLAARKAGSPRSHSTRENC
						RRSRCPDTAHRRRRRGRRRNPSCVRSPRWR
1191	2541	A	9489	1	411	LADALCLSAAATGAVRPGARAQPSTRRRLSP
		,				SVRVCCRAAAASNLLYSSCLQRHSERASEEG
			1			ERGSLSAKCCSLVLRGGCSSSNSIISFRRIT*EI
			1			MAAFVLLSYEQRPLKRPRLGPPDVYPPDPKQ
						KEEELTAVNVK
1192	2542	A	9497	389	161	VSFLSMSSGHCIRSTRGSKMVSWSVIAKIQEI*
Ī .				ĺ		CEEDERKMAREFLAEFMSTYVMMNIHMIVE
						KDTYSDHEEINTS
1193	2543	A	9509	186	1	IAKSQ*KRWQRSGAMETLKHGWWECKLVQF
						FGKTFVNVN*S*TYVYPCDKIILLLGLYPTEM
1194	2544	A	9512	58	433	PLQRSKCLTLRCLRAKPWAWSQSPRACSSAL
				-		LKSSRSRASSLNVQCILQSNPQGHQRI*KQKA
			1	ĺ		SSKGQQFRR*KEHPFMLKTLNKLRIEGT*LKI
						RRAIYDNPTANIIVEGQKLEAFPLRTGTRQ
1195	2545	A	9515	595	1223	GHGAPSFQTQVPRTP*ASWPVVPAASESAPAP
			1			AGGGASLPVAAGSCAAAPHTEPGAPQHLLDC
						PCPLCLARPPRRPLPDTCYGPGSGRSASLAEPP
			1			LPRCSCAPLRSASAPQVS*CV*AVNLLPHNL*
			ľ			PLHLLHD*EKAWGFLFSSASHCFQGQICLLP
						APGSGPCGATARPSRGGRAGGSRARRPIPPGP
						GTRRTPSGCQNPAASGG
	L	L	J	L	L	1

	SEQ ID NO: of	SEQ ID NO: of	Met hod	SEQ ID NO:	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid,
194	eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
minto acid residue of peptide residue of peptide sequence   T=Threonine, V=Valine, W=Thyrlophan, possible melecutide deletion,  =possible melecutide  =possible		uence					
					amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
196		,				sequence	
1996							
197   2547   A   9521   289   448	1196	2546	A	9518		468	
1197							1
HAMBHLITONISMAIRPL#ADSRPQVQRARYE	1107	2547	٨	0521	280	149	
APNWKYKYGY=PVDMLC	1137	2547	A	9521	209	1440	
1199							APNWKYKYGY*IPVDMLC
199	1198	2548	A	9524	204	1	
199							
1200   2550   A   9548   186	1199	2549	A	9546	1785	1943	
1201   2551   A   9549   591   2   SSVVEFPGPRSSLPLDSTPFCGSSPNTGGC   GSCPSGE*LVSPGSEQRKKYSNSNVIMHETSQ   YHVQHLATFIMDKSEATISVIDDAIRRLYQDLS   KEKIWTQEMLLQVNDQSLRLLDIESQEELEDF   PLPTVQRSQTVLNQLRYFSVLLLVCQDSEQSK   PLPTVQRSQTVLNQLRYFSVLLLVCQDSEQSK   PLPTVQRSQTVLNQLRYFSVLLLVCQDSEQSK   PLPTVQRSQTVLNQLRYFSVLLLVCQDSEQSK   PLYTVQRSQTVLNQLRYFSVLLLVCQDSEQSK   PLYTVQRSQTVLNQLRYFSVLLLVCQDSEQSK   PLYTVQRSQTVLNQLRYFSVLLDVCDSEQSK   PLYTVQRSQTVLNQLRYFSVLLDVCDLSEQSK   PLYTVQRSQTVLNQLRYFSVLLDVCDSEQSK   PLYTVQRSQTVLNQLRYFSVLLTQLSEQSELEDF   PLYTVQRSQTVLNQLRYFSVLLTQLSEQSELEDF   PLYTVQRSQTVSQTVSVVVGIDCQV   SKPRATPFLFCSLHTT   STATE   STAT	1000						
1201	1200	2550	A	9548	186	1	
GSCPSGE=LVSPGSEQRKKXYSNSNVIMHETSO	1201	2551	A	9549	591	2	
REKIWTQEMILQVNDQS,ILDIJESQEELEDF							GSCPSGE*LVSPGSEQRKKYSNSNVIMHETSQ
PIPTYORSQIVLNOLRYPSVILLUCODSEQSK   PDVMFFFELDEVEAELVHEYMESALTDCRLGK   AMRP							
PDVHFFICDEVEAÊLVHEYMESALTDCRLĞK	1			É		ĺ	
1202							
LDCERPPGPLPSLPELAKTSYSDLTGLATED							
1203   2553   A   9568   517   738   RRFFERKOKO/*RYREGROYRORDKMKEWG EKERRREKGEREERKMRHRERKGESGORD TMENWRVERLTEKER     1204   2554   A   9573   83   415   EDKRLRLVDGDSRCGRV*HYRDFWGTLOT TMENWRVERLTEKER     1205   2555   A   9577   64   424   ARGSCPTRFFANGRMGETKDAPQMLVTFK DWAVTFFREEWRQLVLVHRTLYR*GMLETC GLLDTLRINVPQDVVHLLYHGTQLLIVKRE VSHSPCAGDMRELFTREATLTPHEYNNGA     1206   2556   A   9584   38   476   TLGAVLFSEVSKESSTSHSGGQLGRQNRHPKL SNFITPSSPRLAPTANSORNLOGLINMFLTAV NPQPLSTPSWQIETKYSTKVLTGNWMEERRK GLPYKHLTHHQEPPHRYLISTYDDHYNRHG YNPGLPPLRTWNGQKLLWL     1207   2557   A   9586   2   412   LRSSPAALLRALCITTVTGTALALRSRVATTN PDGCRNVLRPKYYRLCKAESWGIALETVT GVAVTSWAMMLTVLTLVCKGQDYNRRQKLP THILCLL*EKGIFGLTFAFIIGLDGSTGPTRFFL FGILFSICFS     1208   2558   A   9597   122   3   IKNYWPGMVAHACNPSPLGGRGRWIA*AQK FADAWADAW     1209   2559   A   9611   148   558   KSLRNVWDLLINITWKADRFFCHSSRTSTIRK GDPGPTFSKMSIWTSGRTSSSYRHDEKRNYQ RIRDHDLLDKRKTYALKAGEDRALLGLAM MVCSIMM*FLIGTILLSTRYGVWTRESQCT LLNASITETFNC     1210   2560   A   9618   384   2   SLHDMLMLAEQQQKWAVNTQNTAWSNA DSKFGQRLEEKMEWSKGRGLGGQGGDDDI KVQVKNNDLGLQATNNEANWIAHQDDPNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK	1202	2552	A	9552	428	1	
LI+YRATYSALPNFSQPTQSSQVSVVGIDGQV							
1203	1						
EKEKRRRÈĞEREERKMÈHRERKGESGQRD   TMENWRVERLITEKER	1000	0.550		0560	515	700	
1204	1203	2553	A	9568	517	738	
DGWDLSDAHVVCQKLGCGVAFNATVSAHFG   EGSGPWILDDLNCTGTESHLWQCPSRGWQQ   HDCRHKEDAGVICSEFTALR     1205							
EGSGPIWLDDLNCTGTESHLWQCPSRGWGQ   HDCRHKEDAGVICSETTALR	1204	2554	A	9573	83	415	" "
HDCRHKEDAGVICSEFTALR	1					}	
1205   2555   A   9577   64   424   ARGSCPTRPTANGRMGETKDAPQMLVTFK DVAVTFFREWRQLVLVHRTLYR*GMLETC GLDTLRIHNYDQDVVHILLYHGTQLLIVKRE VSHSPCAGDMRELFTREATLTPHPYNNGA     1206   2556   A   9584   38   476   TLGAVLFSEVSKESSTSHSGGQLGRQNRHPKL SNFITPSSPRLKP*TASSQRNLGQILNMFLTAV NPQPLSTPSWQIETKYSTKVLTGNWMEERRK GLPYKHLITHHQEPPHRYLISTYDDHYNRHG YNPGLPPLRTWNGQKLLWL     1207   2557   A   9586   2   412   LRSSPAALLRALCITTVTGTALALRSRVATTN PDGCRNVLRFKYYRLCDKAESWGIALETVPT GVAVTSWAIMLTVLTLVCKGQDYNRRQKLP THILCLL*EKGIFGLTFAFIIGLDGSTGPTRFFL FGILFSICFS     1208   2558   A   9597   122   3   IKNYWPGMVAHACNPSPLGGRGRWIA*AQK FADAWADAW     1209   2559   A   9611   148   558   KSLRNVWDLLNNTWKADRFFCHSSRTSTIRK GDPGPTFSKMSIWTSGRTSSSYRHDEKRNIYQ RIRDHDLLDKRKTVTALKAGEDRAILLGLAM MVCSIMM*FILGITLLRSYMQSVWTRESQCT LLNASITETFNC     1210   2560   A   9618   384   2   SLHDMLMLAEQQQKQKWAVNTQNTAWSNA DSKFGQRILEKMEWSKGRGLGVQEQGGPDDI KVQVKNNDLGLQATINNEANWIAHQDDFNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK							
Cold Cold Cold Cold Cold Cold Cold Cold	1205	2555	. <b>A</b>	9577	64	424	ARGSCPTRPRTANGRMGETKDAPQMLVTFK
VSHSPCAGDMRELFTREATLTPHPYNNGA							
1206 2556 A 9584 38 476 TLGAVLFSEVSKESSTSHSGGQLGRQNRHPKL SNFITPSSPRLKP*TASSQRNLGQILNMFLTAV NPQPLSTPSWQIETKYSTKVLTGNWMEERRK GLPYKHLITHHQEPPHRYLISTYDDHYNRHG YNPGLPPLRTWNGQKLLWL  1207 2557 A 9586 2 412 LRSSPAALLRALCITTVTGTALALRSRVATTN PDGCRNVLRPKYYRLCDKAESWGIALETVPT GVAVTSWAIMLTVLTLVCKGQDYNRRQKLP THILCLL*EKGIFGLTFAFIIGLDGSTGPTRFFL FGILFSICFS  1208 2558 A 9597 122 3 IKNYWPGMVAHACNPSPLGGRGRWIA*AQK FADAWADAW  1209 2559 A 9611 148 558 KSLRNVWDLLNNTWKADRFFCHSSRTSTIRK GDPGPTFSKMSIWTSGRTSSSYRHDEKRNIYQ RIRDHDLLDKRKTVTALKAGEDRAILLGLAM MVCSIMM*FLLGITLLRSYMQSVWTRESQCT LLNASITETFNC  1210 2560 A 9618 384 2 SLHDMLMLAEQQQKQKWAVNTQNTAWSNA DSKFGQRILEKMEWSKGRGLGVQEQGFDDI KVQVKNNDLGLQATINNEANWIAHQDDFNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK							
NPQPLSTPSWQIETKYSTKVLTGNWMEERRK GLPYKHLITHHQEPPHRYLISTYDDHYNRHG YNPGLPPLRTWNGQKLL WL LRSSPAALLRALCITTVTGTALALRSRVATTN PDGCRNVLRPKYYRLCDKAESWGIALETVPT GVAVTSWAIMLTVLTLVCKGQDYNRRQKLP THILCLL*EKGIFGLTFAFIIGLDGSTGPTRFFL FGILFSICFS	1206	2556	A	9584	38	476	TLGAVLFSEVSKESSTSHSGGQLGRQNRHPKL
GLPYKHLITHHQEPPHRYLISTYDDHYNRHG   YNPGLPPLRTWNGQKLLWL     1207   2557							
YNPGLPPLRTWNGQKLLWL							NPQPLSIPSWQLEIKYSTKVLTGNWMEERRK   GLPYKHLITHHOEPPHRYLISTVDDHVNRHG
1207 2557 A 9586 2 412 LRSSPAALLRALCITTVTGTALALRSRVATTN PDGCRNVLRPKYYRLCDKAESWGIALETVPT GVAVTSWAIMLTVLTLVCKGQDYNRRQKLP THILCLL*EKGIFGLTFAFIIGLDGSTGPTRFFL FGILFSICFS  1208 2558 A 9597 122 3 IKNYWPGMVAHACNPSPLGGRGRWIA*AQK FADAWADAW  1209 2559 A 9611 148 558 KSLRNVWDLLNNTWKADRFFCHSSRTSTIRK GDPGPTFSKMSIWTSGRTSSSYRHDEKRNIYQ RIRDHDLLDKRKTVTALKAGEDRAILLGLAM MVCSIMM*FLLGITLLRSYMQSVWTRESQCT LLNASITETFNC  1210 2560 A 9618 384 2 SLHDMLMLAEQQKQKWAVNTQNTAWSNA DSKFGQRILEKMEWSKGRGLGVQEQGGPDDI KVQVKNNDLGLQATINNEANWIAHQDDFNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK							YNPGLPPLRTWNGQKLLWL
GVAVTSWAIMLTVLTLVCKGQDYNRRQKLP THILCLL*EKGIFGLTFAFIIGLDGSTGPTRFFL FGILFSICFS  1208 2558 A 9597 122 3 IKNYWPGMVAHACNPSPLGGRGRWIA*AQK FADAWADAW  1209 2559 A 9611 148 558 KSLRNVWDLLNNTWKADRFFCHSSRTSTIRK GDPGPTFSKMSIWTSGRTSSSYRHDEKRNIYQ RIRDHDLLDKRKTVTALKAGEDRAILLGLAM MVCSIMM*FLLGITLLRSYMQSVWTRESQCT LLNASITETFNC  1210 2560 A 9618 384 2 SLHDMLMLAEQQKQKWAVNTQNTAWSNA DSKFGQRILEKMEWSKGRGLGVQEQGGPDDI KVQVKNNDLGLQATINNEANWIAHQDDFNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK	1207	2557	A	9586	2	412	LRSSPAALLRALCITTVTGTALALRSRVATTN
THILCLL*EKGIFGLTFAFIIGLDGSTGPTRFFL FGILFSICFS  1208 2558 A 9597 122 3 IKNYWPGMVAHACNPSPLGGRGRWIA*AQK FADAWADAW  1209 2559 A 9611 148 558 KSLRNVWDLLNNTWKADRFFCHSSRTSTIRK GDPGPTFSKMSIWTSGRTSSSYRHDEKRNIYQ RIRDHDLLDKRKTVTALKAGEDRAILLGLAM MVCSIMM*FLLGITLLRSYMQSVWTRESQCT LLNASITETFNC  1210 2560 A 9618 384 2 SLHDMLMLAEQQKQKWAVNTQNTAWSNA DSKFGQRILEKMEWSKGRGLGVQEQGGPDDI KVQVKNNDLGLQATINNEANWIAHQDDFNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK							· · · · · · · · · · · · · · · · · · ·
FGILFSICFS   1208   2558   A   9597   122   3   IKNYWPGMVAHACNPSPLGGRGRWIA*AQK FADAWADAW   1209   2559   A   9611   148   558   KSLRNVWDLLNNTWKADRFFCHSSRTSTIRK GDPGPTFSKMSIWTSGRTSSSYRHDEKRNIYQ RIRDHDLLDKRKTVTALKAGEDRAILLGLAM MVCSIMM*FLLGITLLRSYMQSVWTRESQCT LLNASITETFNC   1210   2560   A   9618   384   2   SLHDMLMLAEQQKQKWAVNTQNTAWSNA DSKFGQRILEKMEWSKGRGLGVQEQGGPDDI KVQVKNNDLGLQATINNEANWIAHQDDFNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK			'				
1209 2559 A 9611 148 558 KSLRNVWDLLNNTWKADRFFCHSSRTSTIRK GDPGPTFSKMSIWTSGRTSSSYRHDEKRNIYQ RIRDHDLLDKRKTVTALKAGEDRAILLGLAM MVCSIMM*FLLGITLLRSYMQSVWTRESQCT LLNASITETFNC  1210 2560 A 9618 384 2 SLHDMLMLAEQQQKQKWAVNTQNTAWSNA DSKFGQRILEKMEWSKGRGLGVQEQGGPDDI KVQVKNNDLGLQATINNEANWIAHQDDFNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK							FGILFSICFS
1209 2559 A 9611 148 558 KSLRNVWDLLNNTWKADRFFCHSSRTSTIRK GDPGPTFSKMSIWTSGRTSSSYRHDEKRNIYQ RIRDHDLLDKRKTVTALKAGEDRAILLGLAM MVCSIMM*FLLGITLLRSYMQSVWTRESQCT LLNASITETFNC  1210 2560 A 9618 384 2 SLHDMLMLAEQQQKQKWAVNTQNTAWSNA DSKFGQRILEKMEWSKGRGLGVQEQGGPDDI KVQVKNNDLGLQATINNEANWIAHQDDFNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK	1208	2558	Α	9597	122	3	
GDPGPTFSKMSIWTSGRTSSSYRHDEKRNIYQ RIRDHDLLDKRKTVTALKAGEDRAILLGLAM MVCSIMM*FLLGITLLRSYMQSVWTRESQCT LLNASITETFNC  1210 2560 A 9618 384 2 SLHDMLMLAEQQQKQKWAVNTQNTAWSNA DSKFGQRILEKMEWSKGRGLGVQEQGGPDDI KVQVKNNDLGLQATINNEANWIAHQDDFNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK	1209	2559	A	9611	148	558	
MVCSIMM*FLLGITLLRSYMQSVWTRESQCT LLNASITETFNC  1210 2560 A 9618 384 2 SLHDMLMLAEQQQKQKWAVNTQNTAWSNA DSKFGQRILEKMEWSKGRGLGVQEQGGPDDI KVQVKNNDLGLQATINNEANWIAHQDDFNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK			••		- 10		
1210 2560 A 9618 384 2 SLHDMLMLAEQQKQKWAVNTQNTAWSNA DSKFGQRILEKMEWSKGRGLGVQEQGGPDDI KVQVKNNDLGLQATINNEANWIAHQDDFNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK							
1210 2560 A 9618 384 2 SLHDMLMLAEQQKQKWAVNTQNTAWSNA DSKFGQRILEKMEWSKGRGLGVQEQGGPDDI KVQVKNNDLGLQATINNEANWIAHQDDFNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK							
DSKFGQRILEKMEWSKGRGLGVQEQGGPDDI KVQVKNNDLGLQATINNEANWIAHQDDFNW LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK	1210	2560	A	9618	384	2	
LLAELNTCQRQETADS***WSPKNSHVGKDS GELSAK					-		DSKFGQRILEKMEWSKGRGLGVQEQGGPDDI
GELSAK							
	1211	2561	A	9620	316	610	

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  LGRAWWLTPVIPTLWEAKAGGSPE*D*AGRG
1212	2562	A	9623	297	344	GSRL*SQHFGRPRRVDHLRSAVQDQPGQHGE TPSLLKIQKIN*VWGRRL*SSYSEAEAGESL QFPVDGDYQKIEKITQLFQAQNLSLCLAMTR
1212	2302		9023	291	344	TREL*KGGGKGRHE*AVVPFLKKGGYGVKAP AILNTSNCT*CF*ETKMLSDDPKACVFEVSSA DL*NTSFGVIR
1213	2563	A	9624	2	356	AELSLASTACGRNTSGDSLPDYDRAPISSPLA TSGTILSAISCLWDLPTPVLRVGLSCQPSMSSQ IPRMYSTDVEAAVNSLEDLYLQAYYAYLCVG LYFHRDDMALEGVSRFL*ELAE
1214	2564	A	9634	776	912	SLSRWVRAKL*VPYNQENCLNPRGGGCSEPR SHYCTPAWATEKDS
1215	2565	A	9636	220	426	KPGNFAVSSEY*DITSGQLKTAVRG*IEMTST EENFGEKLHDIGFGNGFLDKT*KAQATKAKI DK
1216	2566	A	9637	391	76	CFLEDGCTQAS*AEEAAVSPSMAEEEQGSTSC RERRSIRFKMKNHSPDDTIKENVTISNIRTRKI NHLPETERNLLEHGLMYIRLNAAFCSLVAHS LFGFILKAT
1217	2567	A	9655	2008	2432	LHCKMGALETQTHPCSQNMLRSLQKCCCKV EEHHLQPVQVLQTLLHSATAGTGCRRPARPP PAPPTPTPWRSRQSGKQSERAS*LKGRGRYGL GALGGRGGRALGGSRWPPPLPGETLFSGCKH RRRRGSDAAPGEEAGT
1218	2568	A	9658	3	405	HASARALLSPNLSPNNKMAISGGPVLGFFIIA VLMSAQEPWAIKEEHVIIQAEFYLNPDQSGEF MLDFEGEDTFHGDMAKKETVWRLE*LARLD NFEAQRALANIAADQAALEIMDMGSDYTLIP NVPPKVTVL
1219	2569	A	9662	3	284	PDWTEKRKMQDTGSILPLHWFGFGYAALVA YGGIIGYVKAGSVPSLAAGLLFGSLSGLGAYQ LSQDPRNVWVFLATSGTLAGIMGMRFYHSG KL
1220	2570	A	9669	200	699	LLLTGYIQTLQNQQLSGNQQEMQAVDNLTSA PGNTSLCTRDYKITQVLFPLLYTVLFFVGLITN GLAMRIFFQIRSKSNFIIFLKNTVISDLLMILTF PFKILSDAKLGTGPLRTFVCQVTSVIFYFTMYI SISFLGLITIDRYQKTTRPFKTSNPKNLLGAKIL K
1221	2571	A	9676	164	562	KERDSSTFSAAMTTMQGMEQAMPGAGPGVP QLGNMAVIHSHLWKGLQEKFLKGEPKVLGV VQILTALMSLSMGITMMCMASNTYGSNPISV YIGYTIWGSVMFIISGSLSIAAGIRTTKGLVRG SLGMNITSS
1222	2572	A	9688	43	412	VAKMVKCCSAIGCASRCLPNSKLKGLTFHVF PTDENIKRKWVLAMKRLDVNAAGIWEPKKG DVLCSRHFKKTDFDRSAPNIKLKPGVIPSIFDS PYHLQGKREKLHCRKNFTLKTVPATNYNH
1223	2573	A	9696	308	564	RTSMGILYSEPICQAAYQNDFGQVWRWVKE DSSYANVQDGFNGDTPLICACRRGHVRIVSFL LKKECLCQPQKPERENLLALCCE
1224	2574	A	9700	3	632	DAWASGELGSLFDHHVQRAVCDTRAKYRE GRRPRAVKVYTINLESQYLLIQGVPAVGVMK ELVERFALYGAIEQYNALDEYPAEDFTEVYLI KFMNLQSARTAKRKMDEQSFFGGLLHVCYA PEFETVEETRKKLQMRKAYVVKTTENKDHY VTKKKLVTEHKDTEDFRQDFHSEMSGFCKA ALNTSAGNSNPYLPYSCELPLCYFSSK

SEQ ID NO: of nucl- cotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
1225	2575	A	9710	1	163	RSGCVLRMTEWETGAPAVAETPDIKLFGKWS TDDVHINDISLQDYIAGVRLILL
1226	2576	A	9713	82	492	QGLPSFLPAFGPSGSWLGPAPTLGSSCNTVDT ICHGYSEIRPLFYLSFCDLLLGLCWLTETLLYG ASVANKDIICYNLQAVGQIFYISSFLYTVNYI WYLYTELRMKHTQSGQSTSPLVIDYTCRVCQ MAFVFSSLI
1227	2577	A	9720	3	416	GKWKRTQVPLLGEECADMDLARKEFLRGNG LAAGKMNISIDLDTNYAELVLNVGRVTLGEN NRKKMKDCQLRKQQNENVSRAVCALLNSGG GVIKAEVENKGYSYKKDGIGLDLENSFSNML PFVPNFLDFMQNGNYF
1228	2578	A	9723	278	411	EASSSNTVASNVADKTDPHSMNSRVFIGNLN
1229	2579	A	9725	121	902	TLVLQKSDVEAVF  LFAMSGFENLNTDFYQTSYSIDDQSQQSYDY  GGSGGPYSKQYAGYDYSQQGRFVPPDMMQP QQPYTGQIYQPTQAYTPASPQPFYGNNFEDEP PLLEELGINFDHIWQKTLTVLHPLKVADGSIM NETDLAGPMVFCLAFGATLLLAGKIQFGYVY GISAIGCLGMFCLLNLMSMTGVSFGCVASVL GYCLLPMILLSSFAVIFSLQGMVGIILTAGIIG WCSFSASKIFISALAMEGQQLLVAYPCALLYG VFALISVF
1230	2580	A	9739	11	247	TFVLNMNTPKEEFQDWPIVRIAAHLPDLIVYG HFSPERPFMDYFDGVLMFVDISGKCKRDVCL MWMSNRLAWEFTCRA
1231	2581	A	9744	37	1100	TPLFDFWPGFVLSWLQPLSASLRARRAASGPP ACRIMPTTVDDVLEHGGEFHFFQKQMFFLLA LLSATFAPIYVGIVFLGFTPDHRCRSPGVAELS LRCGWSPAEELNYTVPGPGPAGEASPRQCRR YEVDWNQSTFDCVDPLASLDTNRSRLPLGPC RDGWVYETPGSSIVTEFNLVCANSWMLDLFQ SSVNVGFFIGSMSIGYIADRFGRKLCLLTTVLI NAAAGVLMAISPTYTWMLIFRLIQGLVSKAG WLIGYILITEFVGRRYRRTVGIFYQVAYTVGL LVLAGVAYALPHWRWLQFTVALPNFFFLLY YWCIPESPRWLISQNKNAEAMRIIKHIAKKNG KSLPASL
1232	2582	A	9753	164	517	PGPGMQGPPPITPTSWSLPPWRAYVAAAVLC YINLLNYMNWFIIAGVLLDIQEVFQISDNHAG LLQTVFVSCLLLSAPVFGYLGDRHSRKATMS FGILLWSGAGLSSSFISPRYSWLF
1233	2583	A	9757	25	419	LPAPWTERVRKSEGLVGTCLGDPMASPRTVT IVALSVALGLFFVFMGTIKLTPRLSKDAYSEM KRAYKSYVRALPLLKKMGINSILLRKSIGALE VACGIVMTLVPGRPKDVANFFLLLLVLAVLF FHQLV
1234	2584	A	9765	71	456	RLELDWGFSLHFLPVAYLCPLSSGFEMNVQP CSRCGYGVYPAEKISCIDQIWHKACFHCEVC KMMLSVNNFVSHQKKPYCHAHNPKNNTFTS VYHTPLNLNVRTFPEAISGIHDQEDGEQCKSV FHWD
1235	2585	A	9767	52	559	IRSGAMSVDKAELCGSLLTWLQTFHVPSPCA SPQDLSSGLAVAYVLNQIDPSWFNEAWLQGI SEDPGPNWKLKVTSGLLIRGQTGEEMTRDGP ARHMSWVMGRKRDRCLVINHLFIHSSMEYSP CARPGHSARNNTDKNLPHTAIILVTSNTYTTI KINFQAGRSGSCL
1236	2586	Α	9770	352	608	FRGEALTVRFLTKRFIGEYASNFESIYKKHLC

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						LERKQLNLEIYDPCSQTQKAKFSLTSELHWA DGFVIVYDISDRSSFAFAKALI
1237	2587	A	9793	266	515	NILAIIYFPFPRLFLLRDSQSNPKAFALTLCHH QKIKNFQILPVSIDALTPPLVVCFLVSFLTHFS RYKPTRPVCITQFQGCS
1238	2588	A	9802	537	967	ELGAGRSDREAMEAAVKEEISVEDEAVDKNI FRDCNKIAFYRRQKQWLSKKSTYRALLDSVT TDEDSTRFQIINEASKVPLLAEIYGIEGNIFRLK INEETPLKPRFEVPDVLTSKPSTVRLISCSGDT GSLILADGKGDLKC
1239	2589	A	9805	105	540	VPGDPAMVRAGAVGAHLPASGLDIFGDLKK MNKRQLYYQVLNFAMIVSSALMIWKGLIVLT GSESPIVVVLSGSMEPAFHRGDLLFLTNFRED PIRAGEIVVFKVEGRDIPIVHRVIKVHEKDNG DIKFLTKGDNNEGDDRGSYK
1240	2590	A	9819	3	305	TDGRDPLPCAARRRGGGGECCGAGWVAEWS PQPLDPAMLLWMQGFVLEAVACQDNDDYLR YGILFEDLDCNGDGVVDIIELQEGLRNWSSAF DPNSEEHG
1241	2591	A	9834	841	1209	SPARGKSNRTDVMITAPKNKKMTENLAAPEA LDSSTHSSSTATQSRAKMNTPAPTPSTVPAIPR GGSGGPPPCAPHDRVSSVLQCDTQAMDHKTE SSHSVVEFLFKRTKTPSPFHPAVRENRN
1242	2592	A	9843	3	589	TISCGPATEPPASLLSSASSDDFCKEKTEDRYS LGSSLDSGMRTPLCRICFQGPEQGELLSPCRC DGSVKCTHQPCLIKWISERGCWSCELCYYKY HVIAISTKNPLQWQAISLTVIEKVQVAAAILGS LFLIASISWLIWSTFSPSARWQRQDLLFQICYG MYGFMDVMIVAVDSEDMVQAAKEVGKRWS DIPP
1243	2593	A	9846	198	411	WRISHHAGKMPVMKGLLAPQNTFLDTIATRF DGTHSNFILANAQVAKGFPIVYCSDGFCELAG FARTEVMQ
1244	2594	A	9848	116	650	PICGFLYLCSAMASESSPLLAYRLLGEEGVAL PANGAGGPGGASARKLSTFLGVVVPTVLSMF SIVVFLRIGFVVGHAGLLQALAMLLVAYFILA LTVLSVCAIATNGAVQGGGAYCILQHRWTG VWPVLPAREVMISRTLGPEVGGSIGLMFYLA NVCGCAVSLLGLVESVLDVFGA
1245	2595	A	9849	573	1620	KSKCRFPEGLSEGFGPMRKEALSSGSVQEAE AMLDEPQEQAEGSLTVYVISEHSSLLPQDMM SYIGPKRTAVVRGIMHREAFNIIGRRIVQVAQ AMSLTEDVLAAALADHLPEDKWSAEKRRPL KSSLGYEITFSLLNPDPKSHDVYWDIEGAVRR YVQPFLNALGAAGNFSVDSQILYYAMLGVNP RFDSASSSYYLDMHSLPHVINPVESRLGSSAA SLYPVLNFLLYVPELAHSPLYIQDKDGAPVAT NAFHSPRWGGIMVYNVDSKTYNASVLPVRV EVDMVRVMEVFLAQLRLLFGIAQPQLPPKCL LSGPTSEGLMTWELDRLLWARSVENLATATT TLTSLA
1246	2596	A	9850	114	464	PPQLGAQRVREPRHPDVRAPLRVTSPGLRSRS ARSLGRRPRIAMVTVGNYCEAEGPVGPAWM QDGLSPCFFFTLVPSTRMALGTLALVLALPCK RRERPAGADSLSWGAGPRISSYV
1247	2597	A	9851	2	327	FVRNKKMTRSCSAVGCSTRDTVLSRERGLSF HQFPTDTIQRSKWIRAVNRVDPRSKKIWIPGP GAILCSKHFQESDFESYGIRRKLKKGAVPSVS LYKVFKYSSRCTS

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1248	2598	A	9853	58	444	RVDDFVYSKGGKDAGGADVSLACRRQSIPEE FRGITVVELIKKEGSTLGLTISGGTDKDGKPR VSNLRPGGLAARSDLLNIGDYIRSVNGIHLTR LRHDEIITLLKNVGERVVLEVEYELPPPGGCP WT
1249	2599	A	9856	2	1265	LPPPRPSRHRRGRAGTRASAAAAAGPTVSAV RAPVRGQDSGAGTPQGRLAGRGAHLSRVGA SGSGVAAGPAARHAPRRRCADAGEAVGASC GRCAVALLSGVCTLVSTHVCVGSGCPGAAGT PMGAGDAGASAESAVTTAPQEPPARPLQAGS GAGPAPGRAMRSTTLLALLALVLLYLVSGAL VFRALEQPHEQQAQRELGEVREKFLRAHPCV SDQELGLLIKEVADALGGGADPETNSTSNSSH SAWDLGSAFFFSGTIITTIGGGGDWHVGGGK ELPHGGRCRETEGSQVAPRLPASPLCPGYGN VALRTDAGRLFCIFYALVGIPLFGILLAGVGD RLGSSLRHGIGHIEAIFLKWHVPPELVRVLSA MLFLLIGCLLFVLTPTFVFCYMEDWSKLEAIY FVIVTLTTVGFGDYVA
1250	2600	A	9873	2	652	FVVPSPCGGIPGRAPNGASRPTMGNSASRNDF EWVYTDQPHTQRRKEILAKYPAIKALMRPDP RLKWAVLVLVLVQMLACWLVRGLAWRWLL FWAYAFGGCVNHSLTLAIHDISHNAAFGTGR AARNRWLAVFANLPEGVPYAASFKKYHVDH HRYLGGDGLDVDVPTRLEGWFFCTPARKLL WLVLQPFFYSLRPLCVHPKAVTRMEVLNTLV QLA
1251	2601	A	9875	150	1209	PVIMPLHFSPGDIVRPSCCVSSSPKLRRNAHSR LESYRPDTDLSREDTGCNLQHISDRENIDDLN MEFNPSDHPRASTIFLSKSQTDVREKRKSLFIN HHPPGQIARKYSSCSTIFLDDSTVSQPNLKYTI KCVALAIYYHIKNRDPDGRMLLDIFDENLHPL SKSEVPPDYDKHNPEQKQIYRFVRTLFSAAQL TAECAIVTLVYLERLLTYAEIDICPANWKRIV LGAILLASKVWDDQAVWNVDYCQILKDITVE DMNELERQFLELLQFNINVPSSVYAKYYFDL RSLAEANNLSFPLEPLSRERAHKLEAISRLCED KYKDLRRSARKRSASADNLTLPRWSPAIIS
1252	2602	A	9879	6	376	KRPDSRPPAQYRAGPTRPRTRGCELLYWKAT KAVGIKMGSLSTANVEFCLDVFKELNSNNIG DNIFFSSLSLLYALSMVLLGARGETEEQLEKV WNSSEVCSEPRSLSCSRSGSAKLILSLYO
1253	2603	A	9880	180	388	KEQAELLYGLYCQCDLTLSSHPSSVPAMSSC NFTHATFVLIGIPGLEKAHFWVGFPLLSMYVA AMFGNC
1254	2604	A	9881	19	494	VISFQIITDTIMDSSTAHSPVFLVFPPEITASEYE STELSATTFSTQSPLQKLFARKMKILGTIQILF GIMTFSFGVIFLFTLLKPYPRFPFIFLSGYPFWG SVLFINSGAFLIAVKRKTTETLIILSRIMNFLSA LGAIAGIILLTFEFHPRSKLHL
1255	2605	A	9896	72	386	RPGREQRDCFQAPPLGLGGRQTDMMHHPLT GATCVGLPNVGMCPQLSGALTFMYLQQGNQ EATVAPDTMAQPYASAQFAPPQNGIPGEYTA PHPHPAPEYTGOTT
1256	2606	A	9902	95	399	SGGPAGLLHRPVLPKMGLSGLLPILVPFILLG DIQEPGHAEGILGKPCPKIKVECEVEEIDQCTK PRDCPENMKCCPFSRGKKCLDFRKVSLTLYH KEELE
1257	2607	A	9905	374	459	EHLKSTPNRLGVVAHTCNPSTLGGRGGW

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1258	2608	A	9911	364	1974	AGPGVPAVGGRWASGPGLGGRTLCSGPPDH QRRGPSCGASGDPQCVGSPHPQRARPLLARP GARLLPGHLPSPRPPRLPTGQPPAAAFRGPVR PQGGGHIHPLPTPGGRPCFAVSEGSGSALLLS YLGECGSSSYVTGAACISPVLRCREWFEAGLP WPYERGFLLHQKIALSRYATALEDTVDTSRL FRSRSLREFEEALFCHTKSFPISWDAYWDRND PLRDVDEAAVPVLCICSADDPVCGPPDHTILTT ELFHSNPYFFLLLSRHGGHCGFLRQEPLPAWS HEVILESFRALTEFPRTEERIKGLSRHRASFLG GRRRGGALQRREVSSSSNLEEIFNWKRSYTRL MAAAAGAAAAPGSREPQDRPECGAGHPGPR YYRHPERWLLRPEAFLGPLRTRAPSAEDSQR ERPAARSGPEMRVRYPVVAAVLAPYLALSQD PMVKSSASGQGASGSYNHVREEMLIKAGGA MSRRVVRQSKFRHVFGQAAKADQAYEDIRV SKVTWDSSFCAVNPKFLAIIVEAGGGGAFIVL PLAK
1259	2609	A	9919	693	935	GCFKFIGESTCCWIFPSSVTTQCVVAKAPRAA TLSKAERLRSQPGPEQGGSSYRPRTPTAAAIL PPRPGRSHRKRKLVSTK
1260	2610	A	9921	455	1082	QRSCLCSAIEKDGGDVKALYRRSQALEKLGR LDQAVLDLQRCVSLEPKNKVFQEALRNIGGQ IQEKVRYMSSTDAKVEQMFQILLDPEEKGTE KKQKASQNLVVLAREDAGAEKIFRSNGVQLL QRLLDMGETDLMLAALRTLVGICSEHQSRTV ATLSILGTRRVVSILGVESQAVSLAACHLLQV MFDALKEGVKKGFRGKEGAIIV
1261	2611	A	9928	1	438	GFRGAEAPGAAQAPKKKKPRPTEGGPGAGSG RGKDPYRGPTLLHQPKPPKDEFLSSLESYEIAF PTRVDHNGALLAFSPPPPQRQRRGTGATAES RLFYKEASPSTHFLLNLTRSSRLLAGHVSVEY WTREGLAWQRADRPHCLYA
1262	2612	A	9931	168	435	AAEMGRAGAAAVIPGLALLWAVGLGGPPPA PPRLPFCLQELQGRHALHTFSLERTCSYQDFL WADEGRLLHVGAQDLATWHTLSPLGLW
1263	2613	A	9938	247	488	RMSATSVDQRPKGQGNKVSVQNGSIHQKDG CNDDDFEPYLRSPDNQSNSYPPMSDPYMPGY YAPSIGFPYSLGEAAWSQL
1264	2614	A	9941	61	277	ESIGLTALGPRRRPWEHRWSDPITLKMKGWG WLALLLGALLGTAWARRSQDLHCGACKAVR RRVRQFNIYDY
1265	2615	A	9956	2	522	FVASEVSKMPVPASWPHPPGPFLLLTLLLGLT EVAGEEELQMIQPEKLLLVTVGKTATLHCTV TSLLPVGPVLWFRGVGPGRELIYNQKEGHFP RVTTVSDLTKRNNMDFSIRISSITPADVGTYY CVKFRKGSPDHVEFKSGAGTELSVRGEYSVG FLSQVWWWLSSHPFMN
1266	2616	A	10002	243	387	PKNNACHLLFTAVCOPRCKHGECIGPNKCKC HPGYAGKTCNQGRKTV
1267	2617	A	10004	36	707	LPAPASTWSVARETMASSSVPPATVSAATAG PGPGFGFASKTKKKHFVQQKVKVFRAADPLV GVFLWGVAHSINELSQVPPPVMLLPDDFKAS SKIKVNNHLFHRENLPSHFKFKEYCPQVFRNL RDRFGIDDQDYLVSLTRNPPSESEGSDGRFLIS YDRTLVIKEVSSEDIADMHSNLSNYHQVRPLS SPILSLSSLLTYSSAIVSNRCQLGRKLIGRENP
1268	2618	A	10005	2	209	GEGYELFVPSNGVPAVCHMVGRRPHRAVLSP SQDELEHSLGESAAQGAAGVVLWVSWENTR

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seq- uence	uence		09/496 914	correspondi ng to first amino acid residue of peptide sequence	to last amino acid residue of peptide sequence	M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Scrine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide insertion
1269	2619	A	10010	245	688	TKVSLGLA FGMLKNKGHSSKKDNLAVNAVALQDHILHD LQLRNLSVADHSKTQVQKKENKSLKRDTKAI IDTGLKKTTQCPKLEDSEKEYVLDPKPPPLTL AQKLGLIGPPPPPLSSDEWEKVKQRSLLQGDS VQPCPICKEEFELRPQVFSIRG
1270	2620	A	10011	2	588	RVDDFVRPLPPGLMSRSRASIHRGSIPAMSYA PFRDVRGPSTHRTQYVHSPYDRPGWNPRFCII SGNQLLMLDEDEIHPLLIRDRRSESSRNKLLR RTVSVPVEGRPHGEHEYHLGRSRRKSVPGGK QYSMEGAPAAPFRPSQGFLSRRLKSSIKRTKS QPKLDRTSSFRQILPRFRSADHDRYRGWSMW DEIDV
1271	2621	A	10013	209	363	LPAPPNLSPRLSFGFQFPGGNDNYLTITGPSHP FLSGAEVSQSCRRRGGRA
1272	2622	A	10014	7	388	SAVTISWKWRSVMGIQTSPALLASLGAGLVT LLGLAVGSYLVRRSRRPQVTLLDPNEKDLLR LIDKTLSARSPCKHIYLSTRIDGSLSIRPYTPVT SDEDQGYVDIDIKVYLKGVHPTFPEGGKMSH
1273	2623	A	10016	1	3750	MAARTLGRGVGRLLGSLRGLSGQPARPPCGV SAPRRAASGPSGSAPAVAAAAAQPGSYPALS AQAAREPAAFWGPLARDTLVWDTPYHTVW DCDFSTGKIGWFLGGQLNVSVNCLDQHVRKS PESVALIWERDEPGTEVRITYRELLETTCRLA NTLKRHGVHRGDRVAIYMPVSPLAVAAMLA CARIGAVHTVIFAGFSAESLAGRINDAKCKVV ITFNQGLRGGRVVELKKIVDEAVKHCPTVQH VLVAHRTDNKVHMGDLDVPLEQEMAKEDP VCAPESMGSEDMLFMLYTSGSTGMPKGIVHT QAGYLLYAALTHKLVFDHQPGDIFGCVADIG WITGHSYVVYGPLCNGATSVLFESTPVYPNA GRYWETVERLKINQFYGAPTAVRLLLKYGD AWVKKYDRSSLRTLGSVGEPINCEAWEWLH RVVGDSRCTLVDTWWQT
12/4	2024	A	10017		3730	FRPQGTPRSPASHVLTMSAPDEGRRDPPKPKG KTLGSFFGSLPGFSSARNLVANAHSSARARPA ADPTGAPAAEAAQPQAQVAAHPEQTAPWTE KELQPSEKMVSGAKDLVCSKMSRAKDAVSS GVASVVDVAKGVVQGGLDTTRSALTGTKEV VSSGVTGAMDMAKGAVQGGLDTSKAVLTG TKDTVSTGLTGAVNVAKGTVQAGVDTTKTV LTGTKDTVTTGVMGAVNLAKGTVQTGVETS KAVLTGTKDAVSTGLTGAVNVARGSIQTGV DTSKTVLTGTKDTVCSGVTGAMNVAKGTIQT GVDTSKTVLTGTKDTVCSGVTGAMNVAKGT IQTGVDTSKTVLTGTKDTVCSGVTGAMNVA KGTIQTGVDTTKTVLTGTKNTVCSGVTGAVN LAKEAIQGGLDTTKSMVMGTKDTMSTGLTG AANVAKGAMQTGLNTTQNIATGTKDTVCSG VTGAMNLARGTIQTGVDTTKIVLTGTKDTVC SGVTGAANVAKGAVQGGLDTTKSVLTGTKD AVSTGLTGAVNVAKGTVQTGVDTTA KTVLTGTKDTVTTGLVGAVNVAKGTVQTGM DTTKTVLTGTKDTTYTGLVGAVNVAKGTVQTGM DTTKTVLTGTKDTTTGLVGAVNVAKGAVQT GLKTQNIATGTKNTTFGSGVTSAVNVAKGAA QTGVDTAKTVLTGTKDTVTTGLMGAVNVAK GTVQTSVDTTKTVLTGTKDTVCSGVTGAAN

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						VAKGAIQGGLDTTKSVLTGTKDAVSTGLTGA VKLAKGTVQTGMDTTKTVLTGTKDAVCSGV TGAANVAKGAVQMGVDTAKTVLTGTKDTV CSGVTGAANVAKGAVQTGLKTTQNIATGTK NTLGSGVTGAAKVAKGAVQGGLDTTKSVLT GTKDAVSTGLTGAVNLAKGTVQTGVDTSKT VLTGTKDTVCSGVTGAVNVAKGTVQTGVDT AKTVLSGAKDAVTTGVTGAVNVAKGTVQTG VDASKAVLMGTKDTVFSGVTGAMSMAKGA VQGGLDTTKTVLTGTKDAVSAGLMGSGNVA TGATHTGLSTFQNWLPSTPATSWGGLTSSRT TDNGGEQTALSPQEAPFSGISTPPDVLSVGPEP AWEAAATTKGLATDVATFTQGAAPGREDTG LLATTHGPEEAPRLAMLQNELEGLGDIFHPM NAEEQAQLAASQPGPKVLSAEQGSYFVRLGD LGPSFRQRAFEHAVSHLQHGQFQARDTLAQL QDCFRL
1275	2625	A	10025	124	415	TILARKKEKTCPCKKEIGRNSRSGMYSRKAM YKRKYSAANTKVEKKKKEKVLAPVTKPVGG DKNGGTRVVKLPTMPRYYPTEDVPRKLLSHG KKPFS
1276	2626	A	10030	3	507	GGSLRFSPPRVPSCSRVFCPVPPGGCGLPSPMS ASRPQSPTTPWCLPRRYMKHKRDDGPEKQED EAVDVTPVMTCVFVVMCCSMLVLLYYFYDL LVYVVIGIFCLASATGLYSCLAPCVRRLPFGK CRIPNNSLPYFHKRPQARMLLLALFCVAVSV VWGVFRNEDQ
1277	2627	A	10035	51	869	YSRFTVPLPATMASSEVARHLLFQSHMATKT TCMSSQGSDDEQIKRENIRSLTMSGHVGFESL PDQLVNRSIQQGFCFNILCVGETGIGKSTLIDT LFNTNFEDYESSHFCPNVKLKAQTYELQESN VQLKLTIVNTVGFGDQINKEERQLGRSQSTEN PQKYRSEQHPVEPKKCTSFWKGALGKWAGIE SSGQSAQQPYLPINSPPHRLADVADVHLFSSV LSGAFGCYHLDVTVNEFKKQQNRDEQEGYS KGDQEQGSWKHGADPLRGGEM
1278	2628	A	10036	3	457	RAFDVRRKKSLRPCCPRDFHAGCLTVSGPST VMGAVGESLSVQCRYEEKYKTFNKYWCRQP CLPIWHEMVETGGSEGVVRSDQVIITDHPGDL TFTVTLENLTADDAGKYRCGIATILQEDGLSG FLPDPFFQVQVLVSSASSTENSVKTP
1279	2629	A	10039	214	435	NDSLVPMSSWRSCARAPSSESAWRRSAATRR SRKCLRTKRKRWSSGKGTQMQSTLSETPRRA QMPCMWWYPFWG
1280	2630	A	10043	2	344	RATWHNAGKEREAVQLMAGAEKRVKASHS FLRGLFGGNTRIEEACEMYTRAANMFKMAK NWSAAGNAFCQAAKLHMQLQSKHDSATSFV DAGNAYKKADPQGKTARHVACYLCV
1281	2631	A	10080	620	818	VIYKLDSSLFSYFIYFFIFETESHFLPLMKWTG PIMAHCSLKILASRNSADSAFLSAGDTSLSHST
1282	2632	A	10084	3	1640	SASIIIRGDKRASGEVGIAPSSRHILIGEPSAKY NGTAIISLVRGPGILGEVTVFWRIFPPSVGEFA ETSGKLTMRDEQSAVIVVIQALNDDIPEEKSF YEFQLTAVSEGGVLSESSSTANITVVASDSPY GRFAFSHEQLRVSEAQRVNITIIRSSGDFGHVR LWYKTMSGTAEAGLDFVPAAGELLFEAGEM RKSLHVEILDDDYPEGPEEFSLTITKVELQGR GYDFTIQENGLQIDQPPEIGNISIVRIIIMKNDN AEGIIEFDPKYTAFEVEEDVGLIMIPVVRLHGT

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						AVLGRHLVSRIIIAKSDSPFGVIRFLNQSKISIA NPNSTMILSLVLERTGGLLGEIQVNWETVGPN SQEALLPQNRDIADPVSGLFYFGEGEGGVRTII LTIYPHEEIEVEETFIIKLHLVKGEAKLDSRAK DVTLTIQEFGDPNGVVQFAPETLSKKTYSEPL ALEGPLLITFFVRRVKGTFGEIM
1283	2633	A	10088	316	516	MGSKTLPAPVPIHPSLQLTNYSFLQAVNGLPT VPSDHLPNLYGFSALHAVHLHQWTLGYPAM HLXRS
1284	2634	A .	10091	2	569	FVSPSRAMASALIYVSKFKSFVILVVTPLLLLP LVILMPAKFVRCAYVIILMAIYWCTEVIPLAV TSLMPVLLFPLFQILDSRQVCVQYMKDTNML FLGGLIVAVAVERWNLHKRIALRTLLWVGA KPARLMLGFMGVTALLSMWISNTATTAMMV PIVEAILQQMEATSAATEAGLELVDKGKAKE LP
1285	2635	A	10092	290	728	KQSTRPDVMTLYPLHWQEEMSGESVVSSAVP AAATRTTSFKGTSPSSKYVKLNVGGALYYTT MQTLTKQDTMLKAMFSGRMEVLTDSEGWIL IDRCGKHFGTILNYLRDGAVPLPESRREIEELL AEAKYYLVQGLVEECQAALQV
1286	2636	A	10100	1	574	RPRGRGAWAGPGGDYSGVRRQQRRRTRISGS QRGSDAAGTMGCCTGRCSLICLCALQLVSAL ERQIFDFLGFQWAPILGNFLHIIVVILGLFGTIQ YRPRYIMVYTVWTALWVTWNVFIICFYLEVG GLSKDTDLMTFNISVHRSWWREHGPGCVRR VLPPSAHGMMDDYTYVSVTGCIVDFQYLEVI HSA
1287	2637	A	10103	252	376	RSRMGDKPIWEQIGSSFIQHYYQLFDNDRTQL GAIYVSFQL
1288	2638	A	10107	1	478	MEEEDESRGKTEESGEDRGDGPPDRDPTLSPS AFILRAIQQAVGSSLQGDLPNDKDGSRCHGL RWRRCRSPRSEPRSQESGGTDTATVLDMATD SFLAGLVSVLDPPDTWVPSRLDLRPGESEDM LELVAEVRIGDRDPIPLPVPSLLPRLRAWRTG KT
1289	2639	A	10113	237	438	LLSRMPSTNRAGSLKDPEIAELFFKEDPEKLFT DLREIGHGSFGAAYFARDVRTNEVVAIKKMS YSG
1290	2640	A	10114	367	856	RGAKAKSAVLPPGPPCSSILILSPPAPLTPRSPG TEATRPTAMSKSLKKKSHWTSKVHESVIGRN PEGQLGFELKGGAENGQFPYLGEVKPGKVAY ESGSKLVSEELLLEVNETPVAGLTIRDVLAVI KHCKDPLRLKCVKQGESSGLLSVLPGGGTAR GAGQ
1291	2641	A	10116	128	591	RTIRETERRSALSCSVLKSEPLPGLQPQASQQR RRRLPGRRQVQVQEGGGSGLRAWVLAMASV LGSGRGSGGLSSQLKCKSKRRRRRRSKRKDK VSILSTFLAPFKHLSPGITNTEDDDTLSTSSAE VKENRNVGNLAARPPPSGDRARGGATR
1292	2642	A	10121	1	749	QRRRFRAGLWGGHGLTDGLRRNGGCGCSAR VPRVGERLRGHRCPDPLCLLLDMLFLSFHAG SWESWCCCCLIPADRPWDRGQHWQLEMADT RSVHETRFEAAVKVIQSLPKNGSFQPTNEMM LKFYSFYKQATEGPCKLSRPGFWDPIGRYKW DAWSSLGDMTKEEAMIAYVEEMKKIIETMP MTEKVEELLRVIGPFYEIVEDKKSGRSSDITSD

SEQ ID NO: of	SEQ ID NO: of	Met hod	SEQ ID NO:	Predicted beginning	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid,
nucl- eotide	peptide		in USSN	nucleotide location	location	F=Phenylalanine, G=Glycine, H=Histidine,
seq-	seq- uence	ĺ	09/496	correspondi	corresponding to last amino	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline,
uence	donce	ļ	914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
		J	, , ,	amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
				residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
		ł		peptide		/=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion
						LGNVLTSTPNAKTVNGKAESSDSGAESEEEE
1000	2642	<u> </u>	10101			AC
1293	2643	A	10124	2	989	PLMSLVRVVEFVAASSAQKTPSRLENYYMVC
						KADEKFNQLVHFLRNHKQEKHLVFFRYSSGL CGRGIRDSARMCSTCACVEYYGKALEVLVK
1		1	1			GVKIMCIHGKMKYKRNKIFMEFRKLQSGILV
		ļ				CTDVMARGIDIPEVNWVLQYDPPSNASAFVH
						RCGRTARIGHGGSALVFLLPMEESYINFLAIN
-						QKCPLQEMKPQRNTADLLPKLKSMALADRA
1		ļ				VFEKGMKAFVSYVQAYAKHECNLIFRLKDL
						DFASLARGFALLRMPKMPELRGKQFPDFVPV
						DVNTDTIPFKDKIREKQRQKLLEQQRREKTEN
						EGRRKFIKNKAWSKQKAKKK
1294	2644	A	10129	91	1042	VTMYKDCIESTGDYFLLCDAEGPWGIILESLA
1		İ				ILGIVVTILLLLAFLFLMRKIQDCSQWNVLPTQ
						LLFLLSVLGLFGLAFAFIIELNQQTAPVRYFLF GVLFALCFSCLLAHASNLVKLVRGCVSFSWT
						TILCIAIGCSLLQIIIATEYVTLIMTRGMMFVN
1		,		l		MTPCQLNVDFVVLLVYVLFLMALTFFVSKAT
						FCGPCENWKQHGRLIFITVLFSIIIWVVWISML
						LRGNPQFQRQPQWDDPVVCIALVTNAWVFL
		ĺ		l		LLYIVPELCILYRSCRQECPLQGNACPVTAYQ
						HSFQVENQELSRDKWKVLLNSDFLSHSGA
1295	2645	Α	10133	376	518	RPRVVTHNSQWCFLPQDHPGWLPGQSGAPG
1206	0646		10105			GRGAPRQEGPGSSWRQV
1296	2646	A	10135	3	551	EWSLDPFMGIMSGQVGDLSPSQEKSLAQFRE NIQDVLSALPNPDDYFLLRWLQARSFDLQKS
						EDMLRKHMEFRKQQDLANILAWQPPEVVRL
Ì		Ì				YNANGICGHDGEGSPVWYHIVGSQDPKGLLL
ľ						SASKQELLRDSFRSCELLLRECELQSQKLGKR
					:	VEKIIAIFGLEGLGLRDLWKPGIELLQE
1297	2647	A	10138	48	407	MVSSCCGSVCSDQGCGQDLCQETCCRPSCCE
		i			'	TTCCRTTCCRPSCCVSSCCRPQCCQSVCCQPT
						CSRPSCCQTTCCRTTCYRPSCCVSSCCRPQCC
1000	0640		10176			QPVCCQPTCCRPSCCETTCCHPXCC
1298	2648	A	10156	94	453	GGNRKSAEMFSQVPRTPASGCYYLNSMTPEG
						QEMYLRFDQTTRRSPYRMSRILARHQLVTKI QQEIEAKEACDWLRAAGFPQYAQLYEDSQFP
						INIVAVKNDHDFLEKDLGEPLCRRLNT
1299	2649	A	10161	1	393	PRFSELVDGRGRVSARFGGSPSKAATVRSOPT
				_		ASAQLENMEEAPKRVSLALQLPEHGSKDIGN
						VPGNCSENPCQNGGTCVPGADAHSCDCGPGF
		}	}			KGRRCELACIKVSRPCTRLFSETKAFPVWEGG
						VCHHV
1300	2650	A	10162	98	391	AKIASLERIMPANYTCTRPDGDNTDFRYFIYA
						VTYTGILGPGLIGNILALWVFYGYMKETKRA
			,	1		VIFMINLAIADLLQVLSLPLRIFYYLKHDWPF
1301	2651	Ā	10165	1	7545	DCIDVCITSOTCI SONI OENICSVI ABISSIICTE
1301	2031	A	10103	1	1343	PGIRVGITSQTGLSSNLQENCSKLAFISSHGTE KQLQCMPMEGRGRASSSISDLQGKGFEKGTG
						EKHVPGVGSARHSPQASAGGSPWQRGKAQT
1						RWLGKPDPGRKRRRGSPQEEGGLRVSAAAR
			}			LLCSGANRCKVLVRQNSTPNTQQPAVHPSTP
[						PSRPLPQAGRCLVAPLRPHPDWVAAKTLAKA
						LRAPGKPWRLAAPSPLGDLGAPGLPGPSTAP
	' I				•	RTLSVEEPGVECNQLCLYADVTDPVLCLGQK
]						DPGVEGKHCEKEKISSSKELKHVHAKSEPSKP
						ARRLSESLHVVDENKNESKIEREHKRRTSTPV
L			L			IMEGVQEETDTRDVKRQVERSEICTEEPQKQ

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion  KSTLKNEKHLKKDDSETPHLKSLLKKEVKSS  KEKPEREKTPSEDKLSVKHKYKGDCMHKTG
						DETELHSSEKGLKVEENIQKQSQQTKLSSDDK TERKSKHRNERKLSVLGKDGKPVSEYIIKTDE NVRKENNKKERRLSAEKTKAEHKSRRSSDSK
						IQKDSLGSKQHGITLQRRSESYSEDKCDMDST NMDSNLKPEEVVHKEKRRTKSLLEEKLVLKS KSKTQGKQVKVVETELQEGATKQATTPKPD
					 	KEKNTEENDSEKQRKSKVEDKPFEETGVEPV LETASSSAHSTQKDSSHRAKLPLAKEKYKSD
					i ,	KDSTSTRLERKLSDGHKSRSLKHSSKDIKKKD ENKSDDKDGKEVDSSHEKARGNSSLMEKKL SRRLCENRRGSLSQEMAKGEEKLAANTLSTP
		ı			:	SGSSLQRPKKSGDMTLIPEQEPMEIDSEPGVE NVFEVSKTQDNRNNNSHQDIDSENMKQKTS ATVQKDELRTCTADSKATAPAYKPGRGTGV
						NSNSEKHADHRSTLTKKMHIQSAVSKMNPGE KEPIHRGTTEVNIDSETVHRMLLSAPSENDRV QKNLKNTAAEEHVAQGDATLEHSTNLDSSPS
						LSSVTVVPLRESYDPDVIPLFDKRTVLEGSTA STSPADHSALPNQSLTVRESEVLKTSDSKEGG EGFTVDTPAKASITSKRHIPEAHQATLLDGKQ
						GKVIMPLGSKLTGVIVENENITKEGGLVDMA KKENDLNAEPNLKQTIKATVENGKKDGIAVD HVVGLNTEKYAETVKLKHKRSPGKVKDISID
						VERRNENSEVDTSAGSGSAPSVLHQRNGQTE DVATGPRRAEKTSVATSTEGKDKDVTLSPVK AGPATTTSSETRQSEVALPCTSIEADEGLIIGT
						HSRNNPLHVGAEASECTVFAAAEEGGAVVTE GFAESETFLTSTKEGESGECAVAESEDRAADL LAVHAVKIEANVNSVVTEEKDDAVTSAGSEE
						KCDGSLSRDSEIVEGTITFISEVESDGAVTSAG TEIRAGSISSEEVDGSQGNMMRMGPKKETEG TVTCTGAEGRSDNFVICSVTGAGPREERMVT
						GAGVVLGDNDAPPGTSASQEGDGSVNDGTE GESAVTSTGITEDGEGPASCTGSEDSSEGFAIS SESEENGESAMDSTVAKEGTNVPLVAAGPCD
	!	:				DEGIVTSTGAKEEDEEGEDVVTSTGRGNEIGH ASTCTGLGEESEGVLICESAEGDSQIGTVVEH VEAEAGAAIMNANENNVDSMSGTEKGSKDT
				•		DICSSAKGIVESSVTSAVSGKDEVTPVPGGCE GPMTSAASDQSDSQLEKVEDTTISTGLVGGS YDVLVSGEVPECEVAHTSPSEKEDEDIITSVE
						NEECDGLMATTASGDITNQNSLAGGKNQGK VLIISTSTTNDYTPQVSAITDVEGGLSDALRTE ENMEGTRVTTEEFEAPMPSAVSGDDSQLTAS
	ļ					RSEEKDECAMISTSIGEEFELPISSATTIKCAES LQPVAAAVEERATGPVLISTADFEGPMPSAPP EAESPLASTSKEEKDECALISTSIAEECEASVS
					,	GVVVESENERAGTVMEEKDGSGIISTSSVEDC EGPVSSAVPQEEGDPSVTPAEEMGDTAMISTS TSEGCFAVMIGAVLQDEDRI.TITRVEDLSDA
					ļ	AIISTSTAECMPISASIDRHEENQLTADNPEGN GDLSATEVSKHKVPMPSLIAENNCRCPGPVR GGKEPGPVLAVSTEEGHNGPSVHKPSAGOGH
				7.		PSAVCAEKEEKHGKECPEIGPFAGRGQKESTL HLINAEEKNVLLNSLQKEDKSPETGTAGGSST ASYSAGRGLEGNANSPAHLRGPEQTSGQTAK
						DSSVSSIRYLAAVNTGAIKADDMPPVQGTVA EHSFLPAEQQGSEDNLKTSTTKCITGQESKIAP

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide scq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  SHTMIPPATYSVALLAPKCEQDLTIKNDYSGK WTDQASAEKTGDDNSTRKSFPEEGDIMVTVS SEENVCDIGNEESPLNVLGGLKLKANLKMEA YVPSEEEKNGEILAPPESLCGGKPSGIAELQRE PLLVNESLNVENSGFRTNEEIHSESYNKGEISS GRKDNAEAISGHSVEADPKEVEEEERHMPKR KRKQHYLSSEDEPDDNPDVLDSRIETAQRQC PETEPHATKEENSRDLEELPKTSSETNSTTSRV MEEKDEYSSSETTGEKPEQNDDDTIKSQE
1302	2652	A	10167	321	842	EPSLFPFLRPSPARPPPRPPAPFPSPELAGPEPH FVFYFFLSYVHPPKELAKYEYMEEQVILTEKG NSTVAGRGTSVRCLSPSPRPLPPLLPLLADLLE DGFGEHPFYHCLVAEVPKEHWTPEGNPSPFP EARETKCYVRSSVGCVEPLTTQAEVTENLDR KNSQQVFKLLKKK
1303	2653	A	10171	206	429	NMILLKKRRLLINSLGEGTINGLLDELLETNV LSQEDTEIVKCENVTVIDKARDLLDSVIRKGA RACEICITYI
1304	2654	A	10184	970	1524	LCTLSPGISGTAGSCLTTEPGTELGTSFAQNGF YHEAVVLFTQALKLNPQDHRLFGNRSFCHER LGQPAWALADAQVALTLRPGWPRGLFRLGK ALMGLQRFREAAAVFQETLRGGSQPDAAREL RSCLLHLTLQGQRGGICAPPLSPGALQPLPHA ELAPSGLPSLRCPRSTALRSPGLSPLLH
1305	2655	A	10194	2	394	TDLLGRRFRVDGAAMAACEGRRSGALGSSQ SDFLTPPVGGAPWAVATTVVMYPPPPPPPHR DFISVTLSFGESYDNSKSWRRSCWRKWKQL SRLQRNMILFLLAFLLFCGLLFYINLADHWKG IRNTCT
1306	2656	A	10195		410	IPGSTISLEGPLSKWTNVMKGWQYRWFVLDY NAGLLSYYTSKDKMMRGSRRGCVRLRGAVI GIDDEDDSTFTITVDQKTFHFQARDADEREK WIHALEETILRHTLQLQVRVFTWFPDSSLVGA FFFWLVSGFFFK
1307	2657	A	10205	85	308	QGLPSTMVKLGCSFSGKPGKDPGDQDGAAM DSVPLISPLDISQLQPPLPDQVVIKTQTEYQLS SPDQQNYTKSR
1308	2658	A	10214	2	453	ECGGIRQPGPPPPALASAPAATMNRVGGSPS AAANYLLCTNCRKVLRKDKRIRVSQPLTRGP SAFIPEKEVVQANTVDERTNFLVEEYSTSGRL DNITQVMSLHTQYLESFLRSQFYMLRMDGPL PLPYRHYIAIMAAARHQCSYLINM
1309	2659	A	10233	45	421	RGWPEQQSTGRPRDVARQPRCQKEEGRRLRP RALESRTFQGSERSRWGPPLESTKENVQCGH RPAFPNSSWLPFHERLQVQNGECPWQVSIQM SRKHLCGGSILHWWWVLTAAHCFRRTLLDM AV
1310	2660	A	10241	243	442	AFQLFNAKCESAFLSKRNPLQRNWTVLYRRK HKKGQSAEIQKKRTRRAFKFQRAITGASLADI MAK
1311	2661	A	10261	751	176	LPGADYGGGHLSLRLFHLLLTSAAWVPDESQ VTLNSAICVLSTVLIMEFPDLGKHCSEKTCKQ LDFLPVKCDACKQDFCKDHFPYAAHKCPFAF QKDVHVPVCPLCNTPIPVKKGQIPDVVVGDHI DRDCDSHPGKKKEKIFTYRCSKEGCKKKEML QMVCAQCHGNFCIQHRHPLDHSCRHGSRPTI KAG
1312	2662	A	10270	3	669	STSSDEGSPSASTPMINKTGFKFSAEKPVIEVP SMTILDKKDGEQAKALFEKVRKFRAHVEDSD

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion  LIYKLYVVQTVIKTAKFIFILCYTANFVNAISF EHVCKPKVEHLIGYEVFECTHNMAYMLKKL LISYISIICVYGFICLYTLFWLFRIPLKEYSFEKV REESSFSDIPDVKNDFAFLLHMVDQYDQLYS
1313	2663	A	10287	1221	266	KRFGVFLSEVSENKLREISLNHEWTFEKL GAHRVLSPAQGAQPRLRSAASVEVSMVGQR VLLLVAFLLSGVLLSEAAKILTISTLGGSHYLL LDRVSQILQEHGHNVTMLHQSGKFLIPDIKEE EKSYQVIRWFSPEDHQKRIKKHFDSYIETALD GRKESEALVKLMEIFGTQCSYLLSRKDIMDSL KNENYDLVFVEAFDFCSFLIAEKLVKPFVAIL PTTFGSLDFGLPSPLSYVPVFPSLLTDHMDFW GRVKNFLMFFSFSRSQWDMQSTFDNTIKEHF PEGSRPVLSHLLLKAELWFVNSDCAFDFARPL LPNTVYIGGLMEKPIKPVPQVSEPSAFSLGFT
1314	2664	A	10288	536	1890	NVQLAKFSSTLVFFFSCDADPSALAKYVLAL VKKDKSEKELKALCIDQLDVFLQKETQIFVEK LFDAVNTKSYLPPPEQPSSGSLKVEFFPPQEK DIKKEEITKEEEREKKFSRRLNHSPPQSSSRYR ENRSRDERKKDDRSRKRDYDRNPPRRDSYRD RYNRRGRSRSYSRSRSWSKERLRERDRD RSRTRSRSRTRSRERDLVKPKYDLDRTDPLEN NYTPVSSVPSISSGHYPVPTLSSTITVIAPTHHG NNTTESWSEFHEDQVDHNSYVRPMPKKRC RDYDEKGFCMRGDMCPFDHGSDPVVVEDVN LPGMQPFPAQPPVVEGPPPPGLPPPPILTPPPV NLRPPVPPPGPLPPSLPPVTGPPPPLPLQPSG MDAPPNSATSSVPTVVTTGIHHQPPPAPPSLFT ADTYDTDGYNPEAPSITNTSRPMYRHRVHPR AKLG
1315	2665	A	10293	447	1331	SHPLLSCPEKVSAKLRAAAEAAAEERRTRGA GSRGICAGLRSVAPGPEPLKQEEGRREWGSSI GTPSPCGSAQAAAAAAAEEATEKIPALRPALL WALLALWLCCATPAHALQCRDGYEPCVNEG MCVTYHNGTGYCKCPEGFLGEYCQHRDPCE KNRCQNGGTCVAQAMLGKATCRCASGFTGE DCQYSTSHPCFVSRPCLNGGTCHMLSRDTYE CTCQVGFTGRNPKCPGGNLNYQFNGIIVVYS GGSVPPSGTKTSKPAEHNAMGTGSKNFASGT LWVMVSGATSTSTSTL
1316	2666	A	10294	118	572	SLSMESNHKSGDGLSGTQKEAALRALVQRTG YSLVQENGQRKYGGPPPGWDAAPPERGCEIFI GKLPRDLFEDELIPLCEKIGKIYEMRMMMDF NGNNRGYAFVTFSNKVEAKNAIKQLNNYEIR NGRLLGVCASVDNCRLFVGGIPKTKK
1317	2667	A	10301	158	1956	LLKSCGVLLSGVCIPCEGKGPTVLVIQTAVPQ DRPTKSSMRSAAKPWNPAIRAGGHGPDRVRP LPAASSGMKSSKSSTSLAFESRLSRLKRASSE DTLNKPGSTAASGVVRLKKTATAGAISELTES RLRSGTGAFTTTKRTGIPAPREFSVTVSRERSV PRGPSNPRKSVSSPTSSNTPTPTKHLRTPSTKP KQENEGGEKAALESQVRELLAEAKAKDSEIN RLRSELKKYKEKRTLNAEGTDALGPNVDGTS VSPGDTEPMIRALEEKNKNFQKELSDLEEENR VLKEKLIYLEHSPNSEGAASHTGDSSCPTSITQ ESSFGSPTGNQLSSDIDEYKKNIHGNALRTSG SSSSDVTKASLSPDASDFEHITAETPSRPLSSTS NPFKSSKCSTAGSSPNSVSELSLASLTEKIQKM EENHHSTAEELQATLQELSDQQQMVQELTAE

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion  NEKLVDEKTILETSFHQHRERAEQLSQENEKL  MNLLQERVKNEEPTTQEGKIIELEQKCTGILE  QGRFEREKLLNIQQQLTCSLRKVEEENQGAL  EMIKRLKEENEKLNEFLELERHNNNMMAKTL  EECRVTLEGLKMENGSLKSHLQG
1318	2668	A	10303	333	879	GECFIMAAVVQQNDLVFEFASNVMEDERQL GDPAIFPAVIVEHVPGADILNSYAGLACVEEP NDMITESSLDVAEEEIIDDDDDDITLTVEASCH DGDETIETIEAAEALLNMDSPGPMLDEKRINN NIFSSPEDDMVVAPVTHVSVTLDGIPEVMETQ QVQEKYADSPGASSPEQPKRKKK
1319	2669	A	10322	169	654	MEVRMSGSVAVTRAIAVPGLLLLIIATALSL LIGAKSLPASVVLEAFSGTCQSADCTIVLDAR LPRTLAGLLAGGALGLAGALMQTLTRNPLAD PGLLGVNAGASFAIVLGAALFGYSSAQEQLA MAFAGALVASLIVAFTGSQGGGQLSPVRLTL AGVXL
1320	2670	A	10323	441	2	KMNQVAVVIGGGQTLGAFLCHGLAAEGYRV AVVDIQSDKAANVAQEINAEYGESMAYGFG ADATSEQSVLALSRGVDEIFGRVDLLVYSAGI AKAAFISDFQLGDFDRSLQVNLVGYFLCARE FSRLMIRDGIQGRIIQINSKSDE
1321	2671	A	10332	1	453	RHRTAGPGSTISSRTDSASAPAARAMPCEYTY AKLTSDCSRPSLQWYTRAQSKMRRPRLLLKD ILKCTLLVFGVRILYILKLNYTTEECDMKNMH YVDPDHVKRAQKYAQQVLQKESPPKFAKTS MALLFEHRYSVDLLPFVQKAPTDSEA
1322	2672	A	10333	25	423	EPSNGPVVYSALGNEDDEILLLGKDIIGTFAAS ERKMRAHQVLTFLLLFVITSGASENASTSRGC GLDLLPQNVYLCDLDAIWGIVVEAVAGAGA LITLLMLILLGRLPFIKEKEKKSPAVLHFLFL LGTLG
1323	2673	A	10334	52	932	SSLGNEDDEILSLAKDITGMFVASHRKMRAH QVLTFLLLFVITSVASENASTSRGCGLDLLPQ YVSLCDLDAIWGIVVEAAAGAGALITLLLMLI LLVRLPFFKEKEKKSPVGLHFLFLLGTLGP ERLCFPCMQSKIYSYMSPNKCSGMRFPLQEE
		,				NSVTHHEVKCQGKPLAGIYRKREEKRNAGN AVRSAMKSEEQKIKDARKGPLVPFPNQKSEA AEPPKTPPSSCDSTNAAIAKQALKKPIKGKQA PRKKAQGKTQQNRKLTDFYPVRRSSRKSKAE LQSEERKRIDELIESGKEEGMKIDLIDGKGRG VIATKQFSRGDFVVEYHGDLIEITDAKKREAL YAQDPSTGCYMYYFQYLSKTYCVDATRETN RLGRLINHSKCGNCQTKLHDIDGVPHLILIAS RDIAAGEELLYDYGDRSKASIEAHPWLKH
1325	2675	Α	10338	3	870	PGSTISCSELKGTQCRATAGSRGRRPPMTCWL RGVTATFGRPAEWPGYLSHLCGRSAAMDLG PMRKSYRGDREAFEETHLTSLDPVKQFAAWF EEAVQCPDIGEANAMCLATCTRDGKPSARML LLKGFGKDGFRFFTNFESRKGKELDSNPFASL VFYWEPLNRQVRVEGPVKKLPEEEAECYFHS RPKSSQIGAVVSHQSSVIPDREYLRKKNEELE QLYQDQEVPKPKSWGGYVLYPQVMEFWQG QTNRLHDRIVFRRGLPTGDSPLGPMTHRGEE DWLYERLAP
1326	2676	A	10344	2	984	ARAAAHCGICRLVRWWRKRRSVMGIQTSPV LLASLGVGLVTLLGLAVGSYLVRRSRRPQVT LLDPNEKYLLRLLDKTTVSHNTKRFRFALPTA

SEQ ID NO: of	SEQ ID NO: of	Met hod	SEQ ID NO:	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid,
nucl- eotide seq-	peptide seq- uence		in USSN 09/496	nucleotide location correspondi	location corresponding to last amino	F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline,
uence	defice		914	ng to first amino acid	acid residue of peptide	Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan,
		j	<u> </u>	residue of peptide	sequence	Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion  HHTLGLPVGKHIYLSTRIDGSLVIRPYTPVTSD EDQGYVDLVIKVYLKGVHPKFPEGGKMSQY
				3 ;	}	LDSLKVGDVVEFRGPSGLLTYTGKGHFNIQP NKKSPPEPRVAKKLGMIAGGTGITPMLQLIRA
						ILKVPEDPTQCFLLFANQTEKDIILREDLEELQ ARYPNRFKLWFTLDHPPKDWAYSKGFVTAD MIREHLPAPGDDVLVLLCGPPPMVQLACHPN
1327	2677	A	10345	1	968	LDKLGYSQKMRFTY LQSAGEGVTHVLILLESPARPVAAVTQVQRR RYHRLSDMSMLAERRRKQKWAVDPQNTAW
		Transition in the second				SNDDSKFGQRMLEKMGWSKGKGLGAQEQG ATDHIKVQVKNNHLGLGATINNEDNWIAHQ
						DDFNQLLAELNTCHGQETTDSSDKKEKKSFS LEEKSKISKNRVHYMKFTKGKDLSSRSKTDL DCIFGKRQSKKTPEGDASPSTPEENETTTTSAF
}						TIQEYFAKRMAALKNKPQVPVPGSDISETQVE RKRGKKRNKEATGKDVESYLQPKAKRHTEG
1328	2678	A	10346	173	439	KPERAEAQERVAKKKSAPAEEQLRGPCWDQ SSKASAQDAGDHVQPA GSAAMKVKIKCWNGVATWLWVANDENCGI
1329	2679		10351	3	064	CRMAFNGCCPDCKVPGDDCPLVWGQCSHCF HMHCILKWLHAQQVQQHCPMCRQEWKFKE
1329	2079	A	10331	3	964	QMEPGNDTQISEFLLLGFSQEPGLQPFLFGLFL SMYLVTVLGNLLIILATISDSHLHTPMYFFLSN LSFADICVTSTTIPKMLMNIQTQNKVITYIACL
		}				MQMYFFILFAGFENFLLSVMAYDRFVAICHP LHYMVIMNPHLCGLLVLASWTMSALYSLLQI LMVVRLSFCTALEIPHFFCELNQVIQLACSDSF
		<u>.</u>				LNHMVIYFTVALLGGGPLTGILYSYSKIISSIH AISSAQGKYKAFSTCASHLSVVSLFYGAILGV
1330	2680	A	10352	34	2573	YLSSAATRNSHSSATASVMYTVVTPMLNPFI YSLRNKDIKRALGIHLLWGTMKGQFFKKCP IPFLKSCCCCCLFDFPPPPLDQVQEEECEVERV
						TEHGTPKPFRKFDSVAFGESQSEDEQFENDLE TDPPNWQQLVSREVLLGLKPCEIKRQEVINEL
					:	FYTERAHVRTLKVLDQVFYQRVSREGILSPSE LRKIFSNLEDILQLHIGLNEQMKAVRKRNETS VIDQIGEDLLTWFSGPGEEKLKHAAATFCSNQ
						PFALEMIKSRQKKDSRFQTFVQDAESNPLCRR LQLKDIIPTQMQRLTKYPLLLDNIATYTEWPT EREKVKKAADHCRQILNYVNQAVKEAENKQ
						RLEDYQRRLDTSSLKLSEYPNVEELRNLDLTK RKMIHEGPLVWKVNRDKTIDLYTLLLEDILV
	i		<u> </u>			LLQKQDDRLVLRCHSKILASTADSKHTFSPVI KLSTVLVRQVATDNKALFVISMSDNGAQIYE LVAQTVSEKTVWQDLICRMAASVKEQSTKPI
						PLPQSTPGEGDNDEEDPSKLKEEQHGISVTGL QSPDRDLGLESTLISSKPQSHSLSTSGKSEVRD
						LFVAERQFAKEQHTDGTLKEVGEDYQIAIPDS HLPVSEERWALDALRNLGLLKQLLVQQLGLT EKSVQEDWQHFPRYRTASQGPQTDSVIQNSE
						NIKAYHSGEGHMPFRTGTGDIATCYSPRTSTE SFAPRDSVGLAPQDSQASNILVMDHMIMTPE MPTMEPEGGLDDSGEHFFDAREAHSDENPSE
	ļ					GDGAVNKEEKDVNLRISGNYLILDGYDPVQE SSTDEEVASSLTLQPMTGIPAVESTHQQQHSP
						QNTHSDGAISPFTPEFLVQQRWGAMEYSCFEI QSPSSCADSQSQIMEYIHKIEADLEHLKKVEE SYTILCQRLAGSALTDKHSDKS

SEQ ID NO: of nucl- eotide seq- uence	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location correspondi ng to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
1331	2681	A	10353	1	2100	AVEFAEGALTMAPWPELGDAQPNPDKYLEG AAGQQPTAPDKSKETNKTDNTEAPVTKIELLP SYSTATLIDEPTEVDDPWNLPTLQDSGIKWSE RDTKGKILCFFQGIGRLILLLGFLYFFVCSLDIL SSAFQLVGGKMAGQFFSNSSIMSNPLLGLVIG VLVTVLVQSSSTSTSIVVSMVSSSLLTVRAAIP IIMGANIGTSITNTIVALMQVGDRSEFRRAFA GATVHDFFNWLSVLVLLPVEVATHYLEIITQL IVESFHFKNGEDAPDLLKVITKPFTKLIVQLDK KVISQIAMNDEKAKNKSLVKIWCKTFTNKITQ INVTVPSTANCTSPSLCWTDGIQNWTMKNVT YKENIAKCQHIFVNFHLPDLAVGTILLILSLLV LCGCLIMIVKILGSVLKGQVATVIKKTINTDFP FPFAWLTGYLAILVGAGMTFIVQSSSVFTSAL TPLIGIGVITIERAYPLTLGSNIGTTTTAILAAL ASPGNALRSSLQIALCHFFFNISGILLWYPIPFT RLPIRMAKGLGNISAKYRWFAVFYLIIFFFLIP LTVFGLSLAGWRVLVGVGVPVVFIIILVLCLR LLQSRCPRVLPKKLQNWNFLPLWMRSLKPW DAVVSKFTGCFQMRCCCCCRVCCRACCLLC GCPKCCRCSKCCEDLEEAQEGQDVPVKAPET FDNITISREAQGEVPASDSKTECTAL
1332	2682	A	10354	30	1377	SQQGSQPHRQGPPSLLTAPHSLDLPALPPGPR GSQGKLRRVLVPMSVKPSWGPGPSEGVTAVP TSDLGEIHNWTELLDLFNHTLSECHVELSQST KRVVLFALYLAMFVVGLVENLLVICVNWRG SGRAGLMNLYILNMAIADLGIVLSLPVWMLE VTLDYTWLWGSFSCRFTHYFYFVNMYSSIFF LVCLSVDRYVTLTSASPSWQRYQHRVRRAM CAGIWVLSAIIPLPEVVHIQLVEGPEPMCLFM APFETYSTWALAVALSTTILGFLLPFPLITVFN VLTACRLRQPGQPKSRRHCLLLCAYVAVFV MCWLPYHVTLLLLTLHGTHISLHCHLVHLLY FFYDVIDCFSMLHCVINPILYNFLSPHFRGRLL NAVVHYLPKDQTKAGTCASSSSCSTQHSIIIT KGDSQPAAAAPHPEPSLSFQAHHLLPNTSPISP TQPLTPS
1333	2683	Α	10358	2	884	AAGAGADGREPASERASRAEPPAVAMGQND LMGTAEDFADQFLRVTKQYLPHVARLCLIST FLEDGIRMWFQWSEQRDYIDTTWNCGYLLA SSFVFLNLLGQLTGCVLVLSRNFVQYACFGLF GIIALQTIAYSILWDLKFLMRNLALGGGLLLL LAESRSEGKSMFAGVPTMRESSPKQYMQLGG RVLLVLMFMTLLHFDASFFSIVQNIVGTALMI LVAIGFKTKLAALTLVVWLFAINVYFNAFWT IPVYKPMHDFLKYDFFQTMSVIGGLLLVVAL GPGGVSMDEKKKEW
1334	2684	A	10367	59	1562	QAWSLQVALSPFFFPASPSNSFAAAVPQLLFP ELPLPHVPGQESAKRRSARRFLIMSELTKELM ELVWGTKSSPGLSDTIFCRWTQGFVFSESEGS ALEQFEGGPCAVIAPVQAFLLKKLLFSSEKSS WRDCSQEEQKELLCHTLCDILESACCDHSGS YCLVSWLRGKTTEETASISGSPAESSCQVEHS SALAVEELGFERFHALIQKRSFRSLPELKDAV LDQYSMWGNKFGVLLFLYSVLLTKGIENIKN EIEDASEPLIDPVYGHGSQSLINLLLTGHAVSN VWDGDRECSGMKLLGIHEQAAVGFLTLMEA LRYCKVGSYLKISKIPYLDCLASETHLTVFFA KDMALVAPEAPSEQARRVFQTYDPEDNGFIP DSLLEDVMKALDLVSDPEYINLMKNKLDPEG

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1335	2685	A	10375	82	2929	DTPIKRCLQTKWPYIELLWTTDRSPSLN TRTKRRLGREKAMASPPRGWGCGELLLPFML LGTLCEPGSGQIRYSMPEELDKGSFVGNIAKD LGLEPQELAERGVRIVSRGRTQLFALNPRSGS LVTAGRIDREELCAQSPLCVVNFNILVENKM KIYGVEVEIIDINDNFPRFDEELKVKVNENA AAGTRLVLPFARDADVGVNSLRSYQLSSNLH FSLDVVSGTDGQKYPELVLEQPLDREKETVH DLLLTALDGGDPVLSGTTHIRVTVLDANDNA PLFTPSEYSVSVPENIPVGTRLLMLTATDPDE GINGKLTYSFRNEEEKISETFQLDSNLGEISTL QSLDYEESRFYLMEVVAQDGGALVASAKVV VTVQDVNDNAPEVILTSLTSSISEDCLPGTVIA LFSVHDGDSGENGEIACSIPRNLPFKLEKSVD NYYHLLTTRDLDREETSDYNITLTVMDHGTP PLSTESHIPLKVADVNDNPPNFPQASYSTSVT ENNPRGVSIFSVTAHDPDSGDNARVTYSLAE DTFQGAPLSSYVSINSDTGVLYALRSFDYEQL RDLQLWVTASDSGNPPLSSNVSLSLFVLDQN DNTPEILYPALPTDGSTGVELAPRSAEPGYLV TKVVAVDKDSGQNAWLSYRLLKASEPGLFA VGLHTGEVRTARALLDRDALKQSLVVAVED HGQPPLSATFTVTVAVADRIPDILADLGSIKTP IDPEDLDLTLYLVVAVAAVSCVFLAFVIVLLV LRLRRWHKSRLLQAEGSRLAGVPASHFVGV DGVRAFLQTYSHEVSLTADSRKSHLIFPQPNY ADTLLSEESCEKSEPLLMSDKVDANKEERRV QQAPPNTDWRFSQAQRPGTSGSQNGDDTGT WPNNQFDTEMI.QAMILASASEAADGSSTLGG GAGTMGLSARYGPQFTLQHVLQGELGSDYR QNVYIPGSNATLTNAAGKRDGKAPAGGNGN KKKSGKKEKK
1336	2686	A	10379	I	557	RPRRRQPSFSCRVLVLEDPPCFRFTNSMNQEK LAKLQAQVRIGGKGTARRKKKVVHRTATAD DKKLQSSLKKLAVNNIAGIEEVNMIKDDGTVI HFNNPKVQASLSANTFAITGHAEAKPITEMLP GILSQLGADSLTSLRKLAEQFPRQVLDSKAPK PEDIDEEDDDVPDLVENFDEASKNEAN
1337	2687	A	10380		1263	IPGSTISWSPAAARGLSVCRCCRLHPASAMDL FGDLPEPERSPRPAAGKEAQKGPLLFDDLPPA SSTDSGSGGPLLFDDLPPASSGDSGSLATSISQ MVKTEGKGAKRKTSEEEKNGSEELVEKKVC KASSVIFGLKGYVAERKGEREEMQDAHVILN DITEECRPPSSLITRVSYFAVFDGHGGIRASKF AAQNLHQNLIRKFPKGDVISVEKTVKRCLLD TFKHTDEEFLKQASSQKPAWKDGSTATCVLA VDNILYIANLGDSRAILCRYNEESQKHAALSL SKEHNPTQYEERMRIQKAGGNVRDGRVLGV LEVSRSIGDGQYKRCGVTSVPDIRRCQLTPND RFILLACDGLFKVFTPEEAVNFILSCLEDEKIQ TREGKSAADARYEAACNRLANKAVQRGSAD NVTVMVVRIGH
1338	2688	A	10385	3	589	GPSQSMAAGELEGGKPLSGLLNALAQDTFHG YPGITEELLRSQLYPEVPPEEFRPFLAKMRGIL KSIASADMDFNQLEAFLTAQTKKQGGITSDQ AAVISKFWKSHKTKIRESLMNQSRWNSGLRG LSWRVDGKSQSRHSAQIHTPVAIIELELGKYG QESEFLCLEFDEVKVNQILKTLSEVEESISTLIS

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1339	2689	A	10386	50	390	LGAMAKHHPDLIFCRKQAGVAIGRLCEKCDG KCVICDSYVRPCTLVRICDECNYGSYQGRCVI CGGPGVSDAYYCKECTIQEKDRDGCPKIVNL GSSKTDLFYERKKYGFKKR
1340	2690	A	10388		3472	SQLRKGASATHSSPSRTDCIAQMMDIYVCLK RPSWMVDNKRMRTASNFQWLLSTFILLYLM NQVNSQKKGAPHDLKCVTNNLQVWNCSWK APSGTGRGTDYEVCIENRSRSCYQLEKTSIKIP ALSHGDYEITINSLHDFGSSTSKFTLNEQNVSL IPDTPEILNLSADFSTSTLYLKWNDRGSVFPHR SNVIWEIKVLRKESMELVKLVTHNTTLNGKD TLHHWSWASDMPLECAIHFVEIRCYIDNLHFS GLEEWSDWSPVKNISWIPDSQTKVFPQDKVIL VGSDITFCCVSQEKVLSALIGHTNCPLIHLDGE NVAIKIRNISVSASSGTNVVFTTEDNIFGTVIF AGYPPDTPQQLNCETHDLKEIICSWNPGRVTA LVGPRATSYTLVESFSGKYVRLKRAEAPTNES YQLLFQMLPNQEIYNFTLNAHNPLGRSQSTIL VNITEKVYPHTPTSFKVKDINSTAVKLSWHLP GNFAKINFLCEIEIKKSNSVQEQRNVTIKGVE NSSYLVALDKLNPYTLYTFRIRCSTETFWKW SKWSNKKQHLTTEASPSKGPDTWREWSSDG KNLIIYWKPLPINEANGKILSYNVSCSSDEETQ SLSEIPDPQHKAEIRLDKNDYIISVVAKNSVGS SPPSKIASMEIPNDDLKIEQVVGMGKGILLTW HYDPNMTCDYVIKWCNSSRSEPCLMDWRKV PSNSTETVIESDEFRPGIRYNFFLYGCRNQGY QLLRSMIGYIEELAPIVAPNFTVEDTSADSILV KWEDIPVEELRGFLRGYLFYFGKGERDTSKM RVLESGRSDIKVKNITDISQKTLRIADLQGKTS YHLVLRAYTDGGVGPEKSMYVVTKENSVGL IIAILIPVAVAVIVGVVTSILCYRKREWIKETFY PDIPNPENCKALQFQKSVCEGSSALKTLEMNP CTPNNVEVLETRSAFPKIEDTEIVSPVAERPEN RSDAKPENHVVESYCPPIIEEEIPNPAADETGG TAQVIYIDVQSMYQPQAKPEEQENDPVGGA GYKPQMHLPINSTVEDIAAEEDLDKTAGYRP QANVNTWNLVSPDSPRSIDSNSEIVSFGSPCSI NSRQFLIPPKDEDSPKSNGGGWSFTNFFQNKP ND
1341	2691	A	10392	1	5057	MLPPKHLSATKPKKSWAPNLYELDSDLTKEP DVIIGEGPTDSEFFHQRFRNLIYVEFVGPRKTL IKLRNLCLDWLQPETRTKEEIIELLVLEQYLTII PEKLKPWVRAKKPENCEKLVTLLENYKEMY QPEGESLHGVLVVSAGLRCPLGLSASTLLTW SGLDNSLSWAAVGMSCVLWDIELHHDFLGV ATKSVSTHAQGDAAQGLGGTIVRMWARDSN LATGVLLDDNNSDVTSDDDMTRNRRESSPPH SVHSFSGDRDWDRRGRSRDTEPRDRWSHTR NPRSRMPPRDLSLPVVAKTSFEMDREDDRDS RAYESRSQDAESYQNVVDLAEDRKPHNTIQD NMENYRKLLSLGVQLAEDDGHSHMTQGHSS RSKRSAYPSTSRGLKTMPEAKKSTHRRGICED ESSHGVIMEKFIKDVSRSSKSGRARESSDRSQ RFPRMSDDNWKDISLNKRESVIQQRVYEGNA FRGGFRFNSTLVSRKRVLERKRRYHFDTDGK GSIHDQKGCPRKKPFECGSEMRKAMSVSSLS SLSSPSFTESQPIDFGAMPYVCDECGRSFSVIS EFVEHQIMHTRENLYEYGESFIHSVAVSEVQK

SEQ ID NO: of nucl- cotide	SEQ ID NO: of peptide seq- uence	Met hod	SEQ ID NO: in USSN 09/496	Predicted beginning nucleotide location correspondi	Predicted end nucleotide location corresponding to last amino	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline,
seq- uence	dence		914	ng to first amino acid residue of peptide sequence	acid residue of peptide sequence	Q=Glutamine, N=Asparagne, P=Proline, Q=Glutamine, R=Arginine, S=Scrine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
						SQVGGKRFECKDCGETFNKSAALAEHRKIHA RGYLVECKNQECEEAFMPSPTFSELQKIYGK DKFYECRVCKETFLHSSALIEHQKIHFGDDKD NEREHERERERGETFRPSPALNEFQKMYG KEKMYECKVCGETFLHSSSLKEHQKIHTRGN PFENKGKVCEETFIPGQSLKRRQKTYNKEKLC DFTDGRDAFMQSSELSEHQKIHSRKNLFEGR GYEKSVIHSGPFTESQKSHTITRPLESDEDEKA FTISSNPYENQKIPTKENVYEAKSYERSVIHSL ASVEAQKSHSVAGPSKPKVMAESTIQSFDAIN HQRVRAGGNTSEGREYSRSVIHSLVASKPPRS HNGNELVESNEKGESSIYISDLNDKRQKIPAR ENPCEGGSKNRNYEDSVIQSVFFAKPQKSVP GEGSGEFKKDGEFSVPSSNVREYQKARAKKK YIEHRSNETSVIHSLPFGEQTFRPRGMLYECQ ECGECFAHSSDLTEHQKIHDREKPSGSRNYE WSVIRSLAPTDPQTSYAQEQYAKEQARNKCK DFRQFFATSEDLNTNQKIYDQEKSHGEESQGE NTDGEETHSEETHGQETIEDPVIQGSDMEDPQ
1243	2602		10303		1250	KDDPDDKIYECEDCGLGFVDLTDLTDHQKVH SRKCLVDSREYTHSVIHTHSISEYQRDYTGEQ LYECPKCGESFIHSSFLFEHQRIHEQDQLYSM KGCDDGFIALLPMKPRRNRAAERNPALAGSA IRCLLCGQGFIHSSALNEHMRLHREDDLLEQS QMAEEAIIPGLALTEFQRSQTEERLFECAVCG ESFVNPAELADHVTVHKNEPYEYGSSYTHTS FLTEPLKGAIPFYECKDCGKSFIHSTVLTKHKE LHLEEEEEDEAAAAAAAAQEVEANVHVPQ VVLRIQGLNVEAAEPEVEAAEPEVEAAEPEV EAAEPNGEAEGPDGEAAEPIGEAGQPNGEAE QPNGDADEPDGAGIEDPEERAEEPEGKAEEPE GDADEPDGVGIEDPEEGEDQEIQVEEPYYDC HECTETFTSSTAFSEHLKTHASMIIFEPANAFG ECSGYIERASTSTGGANQADEKYFKCDVCGQ LFNDHLSLARHQNTHTG
1342	2692	A	10393	2	1350	GRPRSSSDNRNFLRERAGLSSAAVQTRIGNSA ASRRSPAARPPVPAPPALPRGRPGTEGSTSLS APAVLVVAVAVVVVVSAVAWAMANYIHV PPGSPEVPKLNVTVQDQEEHRCREGALSLLQ HLRPHWDPQEVTLQLFTDGITNKLIGCYVGN TMEDVVLVRIYGNKTELLVDRDEEVKSFRVL QAHGCAPQLYCTFNNGLCYEFIQGEALDPKH VCNPAIFRLIARQLAKIHAIHAHNGWIPKSNL WLKMGKYFSLIPTGFADEDINKRFLSDIPSSQI LQEEMTWMKEILSNLGSPVVLCHNDLLCKNII YNEKQGDVQFIDYEYSGYNYLAYDIGNHFNE FAGVSDVDYSLYPDRELQSQWLRAYLEAYK EFKGFGTEVTEKEVEILFIQVNQFALASHFFW GLWALIQAKYSTIEFDFLGYAIVRFNQYFKM KPEVTALKVPE
1343	2693	Α	10394	102	839	PEAQTSAVLAREKGHLPTMRHEAPMQMASA QDARYGQKDSSDQNFDYMFKLLIIGNSSVGK TSFLFRYADDSFTSAFVSTVGIDFKVKTVFKN EKRIKLQIWDTAGQERYRTITTAYYRGAMGFI LMYDITNEESFNAVQDWSTQIKTYSWDNAQ VILVGNKCDMEDERVISTERGQHLGEQLGFE FFETSAKDNINVKQTFERLVDIICDKMSESLET DPAITAAKQNTRLKETPPPPQPNCAC
1344	2694	A	10395	2	4136	DRPPWNSRVDDFVTNLIHLSSKGHISPAKDTS LQQRTPAEMSPVLHFYVRPSGHEGAASGHTR

SEQ ID NO: of nucl-eotide sequence	SEQ ID NO: of peptide seq- uence	Method	SEQ ID NO: in USSN 09/496 914	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion RKLQGKLPELQGVETELCYNVNWTAEALPSA EETKKLMWLFGCPLLLDDVARESWLLPGSN DLLLEVGPRLNFSTPTSTNIVSVCRATGLGPV DRVETTRRYRLSFAHPPSAEVEAIALATLHDR MTEQHFPHPIQSFSPESMPEPLNGPINILGEGR LALEKANQELGLALDSWDLDFYTKRFQELQR NPSTVEAFDLAQSNSEHSRHWFFKGQLHVDG QKLVHSLFESIMSTQESSNPNNVLKFCDNSSA IQGKEVRFLRPEDPTRPSRFQQQGLRHVVFT AETHNFPTGVCPFSGATTGTGGRIRDVQCTG RGAHVVAGTAGYCFGNLHIPGYNLPWEDLSF QYPGNFARPLEVAIEASNGASDYGNKFGEPV LAGFARSLGLQLPDGQRREWIKPIMFSGGIGS MEADHISKEAPEPGMEVVKVGGPVYRIGVGG GAASSVQVQGDNTSDLDFGAVQRGDPEMEQ KMNRVIRACVEAPKGNPICSLHDQGAGGNG NVLKELSPPAGAIIYTSRFQLGDPTLNALEIW GAEYQESNALLLRSPNRDFLTHVSARECPA CFVGTITGDRRIVLVDDRECPVRRNGQGDAP PTPPPTPVDLELEWVLGKMPRKEFFLQRKPP MLQPLALPPGLSVHQALERVLRPAVASKRY LTNKVDRSVGGLVAQQCVGPLQTPLADVA VVALSHEBLIGAATALGEQPVKSLLDPKVAA RLAVAEALTNLVFALVTDLRDVKCSGNWM WAAKLPGEGAALADACEAMVAVMAALGVA VCPDITATVTPDLKHPEGRGHLLYVALSPGQ HRLGGTALAQCFSQLGEHPPDLDLPENLVRA FSITQGLLKDRLLCSGHDVSDGGLVTCLLEM AFAGNCGLVDLKHYRDAGLHCLELGHTGE AGPHAMVRVSVNGAVVLEEPVGELRALWEE TSFQLDRLQAEPRCVAEERGLRERMGPSYC LPPTFPKASVPREPGGPSPRVAILREGGSNGDR FMAGNCLVLKRYRDAGLHCLELGHTGE AGPHAMVRVSVNGAVVLEEPVGELRALWEE TSFQLDRLQAEPRCVAEERGLRERMGPSYC LPPTFPKASVPREPGFSPRVAILREGGSNGDR FMAGAELRFKRPDTFSLGVCNGCQLLALLG WVGGDPNEDAAEMGPDSQPARPGLLLRHNL SGRYESR WASVRVGPGPSPRVAILREGGSNGDR FMADAFHLAGFEVWDVTMQDLCSGAIGLDT FRGVAFVGGFSYADVLGSAKGWAAAVTFHP RAGAELRFRKRPDTFSLGVCNGCQLLALLG WVGGDPNEDAAEMGPDSQPARPGLLLRHNL SGRYESR WASVRVGPGPALMLRGMEGAVLP VWSAHGEGYVAFSSPELQAQIEARGLAPLHW ADDDGNPTEQYPLNPNGSPGGVAGICSCDGR HLAVMPHPERAVRP WQWAWRPPFDTILTTS PWLQLINARNWTLEGSC GVRGFWAGTMASRAGPRAAGTDGSDFQHRE RVAMHYQMSVTLKYEIKKLIYVHLVIWLLLV AKMSVGHLRLLSHDUGVAM PAMPAGESLAD L
						AKMSVGHLRLLSHDQVAMPYQWEYPYLLSI LPSLLGLLSFPRNNISYLVLSMISMGLFSIAPLI YGSMEMFPAAQQLYRHGKAYRFLFGFSAVSI MYLVLVLAVQVHAWQLYYSKKLLDSWFTST QEKKHK
1346	2696	A	10398	1	718	DDFVRCGPQSAAMGASARLLRAVIMGAPGS GKGTVSSRITTHFELKHLSSGDLLRDNMLRGT EIGVLAKAFIDQGKLIPDDVMTRLALHELKNL TQYSWLLDGFPRTLPQAEALDRAYQIDTVINL NVPFEVIKQRLTARWIHPASGRVYNIEFNPPK TVGIDDLTGEPLIQREDDKPETVIKRLKAYED QTKPVLEYYQKKGVLETFSGTETNKIWPYVY AFLQTKVPQRSQKASVTP
1347	2697	Α	10402	153	1969	KHRQENNALDMAPEIHMTGPMCLIENTNGEL VANPEALKILSAITQPVVVVAIVGLYRTGKSY

SEQ ID	SEQ ID	Met	SEQ	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine,
NO: of	NO: of	hod	ID NO:	beginning	nucleotide	D=Aspartic Acid, E=Glutamic Acid,
nucl-	peptide	1	in	nucleotide	location	F=Phenylalanine, G=Glycine, H=Histidine,
eotide	seq-		USSN	location	corresponding	I=Isoleucine, K=Lysine, L=Leucine,
seq-	uence		09/496	correspondi	to last amino	M=Methionine, N=Asparagine, P=Proline,
uence		ĺ	914	ng to first	acid residue	Q=Glutamine, R=Arginine, S=Serine,
	Ì	}		amino acid	of peptide	T=Threonine, V=Valine, W=Tryptophan,
1	ł	ł		residue of	sequence	Y=Tyrosine, X=Unknown, *=Stop codon,
	1			peptide		/=possible nucleotide deletion, \=possible
				sequence		nucleotide insertion
	1					LMNKLAGKNKGFSLGSTVKSHTKGIWMWCV
1	ł					PHPKKPEHTLVLLDTEGLGDVKKGDNQNDS
		ļ				WIFTLAVLLSSTLVYNSMGTINQQAMDQLYY
1	]	]				VTELTHRIRSKSSPDENENEDSADFVSFFPDFV
						WTLRDFSLDLEADGQPLTPDEYLEYSLKLTQ
	]	}				GTSQKDKNFNLPRLCIRKFFPKKKCFVFDLPI
						HRRKLAQLEKLQDEELDPEFVQQVADFCSYI
	i	ļ				FSNSKTKTLSGGIKVNGPRLESLVLTYINAISR
1	[					GDLPCMENAVLALAQIENSAAVQKAIAHYD
		ļ				QQMGQKVQLPAETLQELLDLHRVSEREATEV
1	[	ĺ				YMKNSFKDVDHLFQKKLAAQLDKKRDDFCK
						QNQEASSDRCSALLQVIFSPLEEEVKAGIYSK
1	ì					PGGYCLFIQKLQDLEKKYYEEPRKGIQAEEIL
						QTYLKSKESVTDAILQTDQILTEKEKEIEVEC
		1				VKAESAQASAKMVEEMQIKYQQMMEEKEKS
	}	)				YQEHVKQLTEKMERERAQLLEEQEKTLTSKL
		!				QEQARVLKERCQGESTQLQNEIQKLQKTLKK
		<u></u>			0.00	KTKRYMSHKLKI
1348	2698	Ā	10404	5	892	TQLPAPLSGVLSRLQLGSGAPLLTWVQETAG
						VAGGAPRRRTPVTMWRLLARASAPLLRVPLS
	•	[				DSWALLPASAGVKTLLPVPSFEDVSIPEKPKL
						RFIERAPLVPKVRREPKNLSDIRGPSTEATEFT
	Ì	Í		}		EGNFAILALGGGYLHWGHFEMMRLTINRSM
						DPKNMFAIWRVPAPFKPITRKSVGHRMGGGK
1	ł	ł		}		GAIDHYVTPVKAGRLVVEMGGRCEFEEVQG
						FLDQVAHKLPFAAKAVSRGTLEKMRKDQEE
1	l					RERNNQNPWTFERIATANMLGIRKVLSPYDL
1240	2600		10100	50	1104	THKGKYWGKFYMPKRV
1349	2699	A	10409	59	1184	LRRNCSALGGLFQTIISDMKGSYPVWEDFINK
						AGKLQSQLRTTVVAAAAFLDAFQKVADMAT
1	}	}				NTRGGTREIGSALTRMCMRHRSIEAKLRQFSS
		1				ALIDCLINPLQEQMEEWKKVANQLDKDHAK EYKKARQEIKKKSSDTLKLQKKAKKGRGDIQ
	]	}				POLDSALODVNDKYLLLEFTEKOAVRKALIE
						ERGRFCTFISMLRPVIEEEISMLGEITHLQTISE
1						DLKSLTMDPHKLPSSSEQVILDLKGSDYSWS
		[				YQTPPSSPSTTMSRKSSVCSSLNSVNSSDSRSS
1	1	(				GSHSHSPSSHYRYRSSNLAQQAPVRLSSVSSH
	1	ļ				DSGFISQDAFQSKSPSPMPPEAPNQRRKEKRE
	1				1	DOGLIGONICONSTOLMELTALINGIAGERICE
	}					PDPNGGGPTTASGPPAAAEEAOPDPSM
1350	2700	Δ	10410	\$11	058	PDPNGGGPTTASGPPAAAEEAQRPRSM
1350	2700	A	10410	511	958	AGRGGPGKPVSWSSGPGSPGQTQRRSWVKST
1350	2700	A	10410	511	958	AGRGGPGKPVSWSSGPGSPGQTQRRSWVKST RGHSSLLPPSQDFVAGLSVILRGTVDDRLNW
1350	2700	A	10410	511	958	AGRGGPGKPVSWSSGPGSPGQTQRRSWVKST RGHSSLLPPSQDFVAGLSVILRGTVDDRLNW AFNLYDLNKDGCITKEEMLDIMKSIYDMMG
1350	2700	A	10410	511	958	AGRGGPGKPVSWSSGPGSPGQTQRRSWVKST RGHSSLLPPSQDFVAGLSVILRGTVDDRLNW

## WHAT IS CLAIMED IS:

1. An isolated polynucleotide comprising a nucleotide sequence selected from the group consisting of SEQ ID NO: 1-1350, a mature protein coding portion of SEQ ID NO: 1-1350, an active domain of SEQ ID NO: 1-1350, and complementary sequences thereof.

- 2. An isolated polynucleotide encoding a polypeptide with biological activity, wherein said polynucleotide hybridizes to the polynucleotide of claim 1 under stringent hybridization conditions.
- 3. An isolated polynucleotide encoding a polypeptide with biological activity, wherein said polynucleotide has greater than about 90% sequence identity with the polynucleotide of claim 1.
- 4. The polynucleotide of claim 1 wherein said polynucleotide is DNA.
- 5. An isolated polynucleotide of claim 1 wherein said polynucleotide comprises the complementary sequences.
- 6. A vector comprising the polynucleotide of claim 1.
- 7. An expression vector comprising the polynucleotide of claim 1.
- 8. A host cell genetically engineered to comprise the polynucleotide of claim 1.
- 9. A host cell genetically engineered to comprise the polynucleotide of claim 1 operatively associated with a regulatory sequence that modulates expression of the polynucleotide in the host cell.
- 10. An isolated polypeptide, wherein the polypeptide is selected from the group consisting of:
  - (a) a polypeptide encoded by any one of the polynucleotides of claim 1; and
  - (b) a polypeptide encoded by a polynucleotide hybridizing under stringent conditions with any one of SEQ ID NO:1-1350.
- 11. A composition comprising the polypeptide of claim 10 and a carrier.
- 12. An antibody directed against the polypeptide of claim 10.

13. A method for detecting the polynucleotide of claim 1 in a sample, comprising:

- a) contacting the sample with a compound that binds to and forms a complex with the polynucleotide of claim 1 for a period sufficient to form the complex; and
- b) detecting the complex, so that if a complex is detected, the polynucleotide of claim 1 is detected.
- 14. A method for detecting the polynucleotide of claim 1 in a sample, comprising:
- a) contacting the sample under stringent hybridization conditions with nucleic acid primers that anneal to the polynucleotide of claim 1 under such conditions;
- b) amplifying a product comprising at least a portion of the polynucleotide of claim 1; and
- c) detecting said product and thereby the polynucleotide of claim 1 in the sample.
- 15. The method of claim 14, wherein the polynucleotide is an RNA molecule and the method further comprises reverse transcribing an annealed RNA molecule into a cDNA polynucleotide.
- 16. A method for detecting the polypeptide of claim 10 in a sample, comprising:
- a) contacting the sample with a compound that binds to and forms a complex with the polypeptide under conditions and for a period sufficient to form the complex; and
- b) detecting formation of the complex, so that if a complex formation is detected, the polypeptide of claim 10 is detected.
- 17. A method for identifying a compound that binds to the polypeptide of claim 10, comprising:
- a) contacting the compound with the polypeptide of claim 10 under conditions sufficient to form a polypeptide/compound complex; and
- b) detecting the complex, so that if the polypeptide/compound complex is detected, a compound that binds to the polypeptide of claim 10 is identified.
- 18. A method for identifying a compound that binds to the polypeptide of claim 10, comprising:

a) contacting the compound with the polypeptide of claim 10, in a cell, under conditions sufficient to form a polypeptide/compound complex, wherein the complex drives expression of a reporter gene sequence in the cell; and

- b) detecting the complex by detecting reporter gene sequence expression, so that if the polypeptide/compound complex is detected, a compound that binds to the polypeptide of claim 10 is identified.
- 19. A method of producing the polypeptide of claim 10, comprising,
- a) culturing a host cell comprising a polynucleotide sequence selected from the group consisting of a polynucleotide sequence of SEQ ID NO: 1-1350, a mature protein coding portion of SEQ ID NO: 1-1350, an active domain of SEQ ID NO: 1-1350, complementary sequences thereof and a polynucleotide sequence hybridizing under stringent conditions to SEQ ID NO: 1-1350, under conditions sufficient to express the polypeptide in said cell; and
  - b) isolating the polypeptide from the cell culture or cells of step (a).
- 20. An isolated polypeptide comprising an amino acid sequence selected from the group consisting of SEQ ID NO: 1351-2700, the mature protein portion thereof, or the active domain thereof.
- 21. The polypeptide of claim 20 wherein the polypeptide is provided on a polypeptide array.
- 22. A collection of polynucleotides, wherein the collection comprises the sequence information of at least one of SEQ ID NO: 1-1350.
- 23. The collection of claim 22, wherein the collection is provided on a nucleic acid array.
- 24. The collection of claim 23, wherein the array detects full-matches to any one of the polynucleotides in the collection.
- 25. The collection of claim 23, wherein the array detects mismatches to any one of the polynucleotides in the collection.
- 26. The collection of claim 22, wherein the collection is provided in a computer-readable format.

27. A method of treatment comprising administering to a mammalian subject in need thereof a therapeutic amount of a composition comprising a polypeptide of claim 10 or 20 and a pharmaceutically acceptable carrier.

28. A method of treatment comprising administering to a mammalian subject in need thereof a therapeutic amount of a composition comprising an antibody that specifically binds to a polypeptide of claim 10 or 20 and a pharmaceutically acceptable carrier.

Pages  $340 \ \rm to \ 1963$  of this application contain amino acid sequence listings. They can be obtained at the address given below.

Les pages 340 to 1963 de cette demande contiennent des listages des séquences d'acides aminés. Elles peuvent être obtenues à l'adresse indiquée ci-dessous.

World Intellectual Property Organization 34, chemin des Colombettes CH-1211 Genève 20